



## INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

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<b>(21) International Application Number:</b> PCT/US97/20201 <b>(22) International Filing Date:</b> 6 November 1997 (06.11.97)  <b>(30) Priority Data:</b> 08/752,307 19 November 1996 (19.11.96) US  <b>(71) Applicant:</b> MILLENNIUM BIOTHERAPEUTICS, INC. [US/US]; 5th floor, 238 Main Street, Cambridge, MA 02142 (US).  <b>(72) Inventors:</b> McCARTHY, Sean, Anthony; Apartment 103, 145 Pinckney Street, Boston, MA 02114 (US). GEARING, David, Paul; 23 Standish Road, Wellesley, MA 02181 (US). LEVINSON, Douglas, Adam; 111 Maple Street, Sherborn, MA 01770 (US).  <b>(74) Agent:</b> MEIKLEJOHN, Anita, L.; Fish & Richardson P.C., 225 Franklin Street, Boston, MA 02110 (US).		<b>(81) Designated States:</b> AU, CA, JP, European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE).  <b>Published</b> <i>With international search report.</i>
<b>(54) Title:</b> METHOD FOR IDENTIFYING GENES ENCODING NOVEL SECRETED OR MEMBRANE-ASSOCIATED PROTEINS		
<b>(57) Abstract</b>		
<p>The invention features a method for identifying a cDNA nucleic acid encoding a mammalian protein having a signal sequence, which method includes the following steps: a) providing library of mammalian cDNA; b) ligating the library of mammalian cDNA to DNA encoding alkaline phosphatase lacking both a signal sequence and a membrane anchor sequence to form ligated DNA; c) transforming bacterial cells with the ligated DNA to create a bacterial cell clone library; d) isolating DNA comprising the mammalian cDNA from at least one clone in the bacterial cell clone library; e) separately transfecting DNA isolated from clones in step (d) into mammalian cells which do not express alkaline phosphatase to create a mammalian cell clone library wherein each clone in the mammalian cell clone library corresponds to a clone in the bacterial cell clone library; f) identifying a clone in the mammalian cell clone library which express alkaline phosphatase; g) identifying the clone in the bacterial cell clone library corresponding to the clone in the mammalian cell clone library identified in step (f); and h) isolating and sequencing a portion of the mammalian cDNA present in the bacterial cell library clone identified in step (g) to identify a mammalian cDNA encoding a mammalian protein having a signal sequence.</p>		

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METHOD FOR IDENTIFYING GENES ENCODING NOVEL  
SECRETED OR MEMBRANE-ASSOCIATED PROTEINS

Background of the Invention

5           The invention relates to methods for identifying genes encoding novel proteins.

          There is considerable medical interest in secreted and membrane-associated mammalian proteins. Many such proteins, for example, cytokines, are important for  
10 inducing the growth or differentiation of cells with which they interact or for triggering one or more specific cellular responses.

          An important goal in the design and development of new therapies is the identification and characterization  
15 of secreted proteins and the genes which encode them. Traditionally, this goal has been pursued by identifying a particular response of a particular cell type and attempting to isolate and purify a secreted protein capable of eliciting the response. This approach is  
20 limited by a number of factors. First, certain secreted proteins will not be identified because the responses they evoke may not be recognizable or measurable. Second, because *in vitro* assays must be used to isolate and purify secreted proteins, somewhat artificial systems  
25 must be used. This raises the possibility that certain important secreted proteins will not be identified unless the features of the *in vitro* system (e.g., cell line, culture medium, or growth conditions) accurately reflect the *in vivo* milieu. Third, the complexity of the effects  
30 of secreted proteins on the cells with which they interact vastly complicates the task of isolating important secreted proteins. Any given cell can be simultaneously subject to the effects of two or more secreted proteins. Because any two secreted proteins

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will not have the same effect on a given cell and because the effect of a first secreted protein on a given cell can alter the effect of a second secreted protein on the same cell, it can be difficult to isolate the secreted  
5 protein or proteins responsible for a given physiological response. In addition, certain secreted and membrane-associated proteins may be expressed at levels that are too low to detect by biological assay or protein purification.

10 In another approach, genes encoding secreted proteins have been isolated using DNA probes or PCR oligonucleotides which recognize sequence motifs present in genes encoding known secreted protein. In addition, homology-directed searching of Expressed Sequence Tag  
15 (EST) sequences derived by high-throughput sequencing of specific cDNA libraries has been used to identify genes encoding secreted proteins. These approaches depend for their success on a high degree of similarity between the DNA sequences used as probes and the unknown genes or EST  
20 sequences.

More recently, methods have been developed that permit the identification of cDNAs encoding a signal sequence capable of directing the secretion of a particular protein from certain cell types. Both Honjo,  
25 U.S. Patent No. 5,525,486, and Jacobs, U.S. Patent No. 5,536,637, describe such methods. These methods are said to be capable of identifying secreted proteins.

The demonstrated clinical utility of several secreted proteins in the treatment of human disease, for  
30 example, erythropoietin, granulocyte-macrophage colony stimulating factor (GM-CSF), human growth hormone, and various interleukins, has generated considerable interest in the identification of novel secreted proteins. The method of the invention can be employed as a tool in the  
35 discovery of such novel proteins.

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Summary of the Invention

The invention features a method for isolating cDNAs and identifying encode secreted or membrane-associated (e.g. transmembrane) mammalian proteins. The method of the invention relies upon the observation that the majority of secreted and membrane-associated proteins possess at their amino termini a stretch of hydrophobic amino acid residues referred to as the "signal sequence." The signal sequence directs secreted and membrane-associated proteins to a sub-cellular membrane compartment termed the endoplasmic reticulum, from which these proteins are dispatched for secretion or presentation on the cell surface.

The invention describes a method in which cDNAs that encode signal sequences for secreted or membrane-associated proteins are isolated by virtue of their abilities to direct the export of the reporter protein, alkaline phosphatase (AP), from mammalian cells. The present method has major advantages over other signal peptide trapping approaches. The present method is highly sensitive. This facilitates the isolation of signal peptide associated proteins that may be difficult to isolate with other techniques. Moreover, the present method is amenable to throughput screening techniques and automation. Combined with a novel method for cDNA library construction in which directional random primed cDNA libraries are prepared, the invention comprises a powerful and approach to the large scale isolation of novel secreted proteins.

The invention features a method for identifying a cDNA nucleic acid encoding a mammalian protein having a signal sequence, which method includes the following steps:

a) providing library of mammalian cDNA;

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b) ligating the library of mammalian cDNA to DNA encoding alkaline phosphatase lacking both a signal sequence and a membrane anchor sequence to form ligated DNA;

5 c) transforming bacterial cells with the ligated DNA to create a bacterial cell clone library;

d) isolating DNA comprising the mammalian cDNA from at least one clone in the bacterial cell clone library;

10 e) separately transfecting DNA isolated from clones in step (d) into mammalian cells which do not express alkaline phosphatase to create a mammalian cell clone library wherein each clone in the mammalian cell clone library corresponds to a clone in the bacterial  
15 cell clone library;

f) identifying a clone in the mammalian cell clone library which express alkaline phosphatase;

g) identifying the clone in the bacterial cell clone library corresponding to the clone in the mammalian  
20 cell clone library identified in step (f); and

h) isolating and sequencing a portion of the mammalian cDNA present in the bacterial cell library clone identified in step (g) to identify a mammalian cDNA encoding a mammalian protein having a signal sequence.

25 A cDNA library is a collection of nucleic acid molecules that are a cDNA copy of a sample of mRNA.

In another aspect, the invention features ptrAP3 expression vector.

In another aspect, the invention features a  
30 substantially pure preparation of ethb0018f2 protein. Preferably, the ethb0018f2 protein includes an amino acid sequence substantially identical to the amino acid sequence shown in FIG. 5 (SEQ ID NO: 5); is derived from a mammal, for example, a human.

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The invention also features purified DNA (for example, cDNA) which includes a sequence encoding a ethb0018f2 protein, preferably encoding a human ethb0018f2 protein (for example, the ethb0018f2 protein of FIG. 5; SEQ ID NO:5); a vector and a cell which includes a purified DNA of the invention; and a method of producing a recombinant ethb0018f2 protein involving providing a cell transformed with DNA encoding ethb0018f2 protein positioned for expression in the cell, culturing the transformed cell under conditions for expressing the DNA, and isolating the recombinant ethb0018f2 protein. The invention further features recombinant ethb0018f2 protein produced by such expression of a purified DNA of the invention.

By "ethb0018f2 protein" is meant a polypeptide which has a biological activity possessed by naturally-occurring ethb0018f2 protein. Preferably, such a polypeptide has an amino acid sequence which is at least 85%, preferably 90%, and most preferably 95% or even 99% identical to the amino acid sequence of the ethb0018f2 protein of FIG. 5 (SEQ ID NO: 5).

By "substantially identical" is meant a polypeptide or nucleic acid having a sequence that is at least 85%, preferably 90%, and more preferably 95% or more identical to the sequence of the reference amino acid or nucleic acid sequence. For polypeptides, the length of the reference polypeptide sequence will generally be at least 16 amino acids, preferably at least 20 amino acids, more preferably at least 25 amino acids, and most preferably 35 amino acids. For nucleic acids, the length of the reference nucleic acid sequence will generally be at least 50 nucleotides, preferably at least 60 nucleotides, more preferably at least 75 nucleotides, and most preferably 110 nucleotides.

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Sequence identity can be measured using sequence analysis software (e.g., Sequence Analysis Software Package of the Genetics Computer Group, University of Wisconsin Biotechnology Center, 1710 University Avenue, 5 Madison, WI 53705).

In the case of polypeptide sequences which are less than 100% identical to a reference sequence, the non-identical positions are preferably, but not necessarily, conservative substitutions for the reference 10 sequence. Conservative substitutions typically include substitutions within the following groups: glycine and alanine; valine, isoleucine, and leucine; aspartic acid and glutamic acid; asparagine and glutamine; serine and threonine; lysine and arginine; and phenylalanine and 15 tyrosine.

Where a particular polypeptide is the to have a specific percent identity to a reference polypeptide of a defined length, the percent identity is relative to the reference peptide. Thus, a peptide that is 50% identical 20 to a reference polypeptide that is 100 amino acids long can be a 50 amino acid polypeptide that is completely identical to a 50 amino acid long portion of the reference polypeptide. It might also be a 100 amino acid long polypeptide which is 50% identical to the reference 25 polypeptide over its entire length. Of course, many other polypeptides will meet the same criteria.

By "protein" and "polypeptide" is meant any chain of amino acids, regardless of length or post-translational modification (e.g., glycosylation or 30 phosphorylation).

By "substantially pure" is meant a preparation which is at least 60% by weight (dry weight) the compound of interest, i.e., a ethb0018f2 protein. Preferably the preparation is at least 75%, more preferably at least 35 90%, and most preferably at least 99%, by weight the



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compound of interest. Purity can be measured by any appropriate method, e.g., column chromatography, polyacrylamide gel electrophoresis, or HPLC analysis.

By "purified DNA" is meant DNA that is not  
5 immediately contiguous with both of the coding sequences with which it is immediately contiguous (one on the 5' end and one on the 3' end) in the naturally occurring genome of the organism from which it is derived. The term therefore includes, for example, a recombinant DNA  
10 which is incorporated into a vector; into an autonomously replicating plasmid or virus; or into the genomic DNA of a prokaryote or eukaryote, or which exists as a separate molecule (e.g., a cDNA or a genomic DNA fragment produced by PCR or restriction endonuclease treatment) independent  
15 of other sequences. It also includes a recombinant DNA which is part of a hybrid gene encoding additional polypeptide sequence.

By "substantially identical" is meant an amino acid sequence which differs only by conservative amino  
20 acid substitutions, for example, substitution of one amino acid for another of the same class (e.g., valine for glycine, arginine for lysine, etc.) or by one or more non-conservative substitutions, deletions, or insertions located at positions of the amino acid sequence which do  
25 not destroy the function of the protein (assayed, e.g., as described herein). Preferably, such a sequence is at least 85%, more preferably 90%, and most preferably 95% identical at the amino acid level to the sequence of FIG. 5 (SEQ ID NO: 5). For nucleic acids, the length of  
30 comparison sequences will generally be at least 50 nucleotides, preferably at least 60 nucleotides, more preferably at least 75 nucleotides, and most preferably 110 nucleotides. A "substantially identical" nucleic acid sequence codes for a substantially identical amino  
35 acid sequence as defined above.

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By "transformed cell" is meant a cell into which (or into an ancestor of which) has been introduced, by means of recombinant DNA techniques, a DNA molecule encoding (as used herein) ethb0018f2 protein.

5 By "positioned for expression" is meant that the DNA molecule is positioned adjacent to a DNA sequence which directs transcription and translation of the sequence (i.e., facilitates the production of ethb0018f2 protein).

10 By "purified antibody" is meant antibody which is at least 60%, by weight, free from the proteins and naturally-occurring organic molecules with which it is naturally associated. Preferably, the preparation is at least 75%, more preferably at least 90%, and most  
15 preferably at least 99%, by weight, antibody.

By "specifically binds" is meant an antibody which recognizes and binds ethb0018f2 protein but which does not substantially recognize and bind other molecules in a sample, e.g., a biological sample, which naturally  
20 includes ethb0018f2 protein.

Unless otherwise defined, all technical and scientific terms used herein have the same meaning as commonly understood by one of ordinary skill in the art to which this invention belongs. Although methods and  
25 materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, the preferred methods and materials are described below. All publications, patent applications, patents, and other references mentioned herein are  
30 incorporated by reference in their entirety. In case of conflict, the present specification, including definitions, will control. In addition, the materials, methods, and examples are illustrative only and not intended to be limiting.

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Other features and advantages of the invention will be apparent from the following detailed description, and from the claims.

#### Brief Description of the Drawings

5           Figure 1 is a schematic drawing of a portion of the ptrAP3 vector.

          Figure 2 is a representation of the DNA sequence of the ptrAP3 vector (SEQ ID NO:1). The bold, underlined portion is the small fragment removed prior to cDNA  
10 insertion sequence. The italic, underlined portion is the alkaline phosphatase sequence.

          Figure 3 is a representation of the amino acid sequence of human placental alkaline phosphatase (Accession No. P05187). The underlined portion is the  
15 signal sequence. The bold, underlined portion is the membrane anchor sequence.

          Figure 4 is a representation of the amino acid sequence of the alkaline phosphatase encoded by ptrAP3.

          Figure 5 is a representation of the cDNA and amino  
20 acid sequence of a portion of a novel secreted protein identified using the method described in Example 1.

          Figure 6 is a representation of an alignment of the amino acid sequence of clone ethb0018f2 (referred to here as 8f2) and proteins containing conserved IgG  
25 domains. The proteins are D38492 (neural adhesion molecule f3); P20241EURO (Drosophila Neuroglian); P32004EURA (human neural adhesion molecule L1); P35331G-CA (chick neural adhesion molecule related protein); Q02246XONI (human Axonin 1); U11031 (rat neural adhesion  
30 molecule BIG1); and X65224 (chicken Neurofascin) are depicted. In this figure, conserved motifs within the IgG domain are highlighted in bold.

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Detailed Description

In general terms, the method of the invention entails the following steps:

1. Preparation of a randomly primed cDNA library  
5 using cDNA prepared from mRNA extracted from mammalian cells or tissue. The cDNA is inserted into a mammalian expression vector adjacent to a cDNA encoding placental alkaline phosphatase which lacks a secretory signal.
2. Amplification of the cDNA library in bacteria.
- 10 3. Isolation of the cDNA library.
4. Transfection of the resulting cDNA library into mammalian cells.
5. Assay of supernatants from the transfected mammalian cells for alkaline phosphatase activity.
- 15 6. Isolation and sequencing of plasmid DNA clones registering a positive score in the alkaline phosphatase assay.
7. Isolation of full length cDNA clones of novel proteins having a signal sequence.

20 The mammalian cDNA used to create the cDNA library can be prepared using any known method. Generally, the cDNA is produced from mRNA. The mRNA can be isolated from any desired tissue or cell type. For example, peripheral blood cells, primary cells, tumor cells, or  
25 other cells may be used as a source of mRNA.

The expression vector harboring the modified alkaline phosphatase gene can be any vector suitable for expression of proteins in mammalian cells.

The mammalian cells used in the transfection step  
30 can be any suitable mammalian cells, e.g., CHO cells, mouse L cells, Hela cells, VERO cells, mouse 3T3 cells, and 293 cells.

Described below is a specific example of the method of the invention. Also described below are two

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genes, one known and one novel, identified using this method.

### Example I

#### Step 1 Generation of Mammalian Signal Peptide Trap cDNA

##### 5 Libraries

##### Vector

A cDNA library was prepared using ptrAP3, a mammalian expression vector containing a cDNA encoding human placental alkaline phosphatase (AP) lacking a  
10 signal sequence (FIG. 1 and FIG. 2, SEQ ID NO:1). When ptrAP3 is transfected into a mammalian cell line, such as COS7 cells, AP protein is neither expressed nor secreted since the AP cDNA of ptrAP3 does not encode a  
15 translation initiating methionine, a signal peptide, or a membrane anchor sequence. FIG. 3 (SEQ ID NO:2) provides the amino acid sequence of naturally occurring AP. FIG. 4 (SEQ ID NO:3) provides the amino acid sequence of the form of AP encoded by ptrAP3. However, insertion of a cDNA encoding a signal peptide sequence into ptrAP3 such  
20 that the signal sequence within the cDNA is fused to and in frame with AP, facilitates both the expression and secretion of AP protein upon transfection of the DNA into COS7 cells or other mammalian cells. The presence of AP activity in the supernatants of transfected COS7 cells  
25 therefore indicates the presence of a signal sequence in the cDNA of interest.

##### cDNA Synthesis and Ligation

cDNA for ligation to the ptrAP3 vector was prepared from messenger RNA isolated from human fetal  
30 brain tissue (Clontech, Palo Alto, CA: Catalog #6525-1) by a modification of a commercially available "ZAP cDNA synthesis kit" (Stratagene; La Jolla, CA: Catalog # 200401). Synthesis of cDNA involved the following steps.

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(a) Single stranded cDNA was synthesized from 5  $\mu$ g of human fetal brain messenger RNA using a random hexamer primer incorporating a XhoI restriction site (underlined); 5'-CTGACTCGAGNNNNNN-3' (SEQ ID NO:4). This  
5 represented a deviation from the Stratagene protocol and resulted in a population of randomly primed cDNA molecules. Random priming was employed rather than the oligo d(T) priming method suggested by Stratagene in order to generate short cDNA fragments, some of which  
10 would be expected to be mRNAs that encode signal sequences.

(b) The single stranded cDNA generated in step (a) was rendered double stranded, and DNA linkers containing a free EcoRI overhang were ligated to both ends of the  
15 double stranded cDNAs using reagents and protocols from the Stratagene ZAP cDNA synthesis kit according to the manufacturer's instructions.

(c) The linker-adapted double-stranded cDNA generated in step (b) was digested with XhoI to generate  
20 a free XhoI overhang at the 3' end of the cDNAs using reagents from the Stratagene ZAP cDNA synthesis kit according to the manufacturers instructions.

(d) Linker-adapted double-stranded cDNAs were size selected by gel filtration through SEPHACRYL™ S-500 cDNA  
25 Size Fractionation Columns (Gibco BRL; Bethesda, MD: Catalog #18092-015) according to the manufacturers instructions.

(e) Size selected, double-stranded cDNAs containing a free EcoRI overhang at the 5' end and a free  
30 XhoI overhang at the 3' end were ligated to the ptrAP3 backbone which had been digested with EcoRI and XhoI and purified from the small, released fragment by agarose gel electrophoresis.

(f) Ligated plasmid DNAs were transformed into E.  
35 Coli strain DH10b by electroporation.

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This process resulted in a library of cDNA clones composed of several million random primed cDNAs (some of which will encode signal sequences) prepared from human fetal brain messenger RNA, fused to the AP reporter cDNA, in the mammalian expression vector ptrAP3.

### Step 2 Plating and Automated Picking of Bacterial Colonies

Next, the transformed bacterial cells were plated, and individual clones were identified. A sample of transformed E. coli containing the random primed human fetal brain cDNA library described in Step 1 was plated for growth as individual colonies, using standard procedures. Each E. coli colony contained an individual cDNA clone fused to the AP reporter in the ptrAP3 expression vector. Approximately 20,000 such E. coli colonies were plated, representing approximately 0.5% of the total cDNA library.

Next, E. coli colonies were picked from the plates and inoculated into deep well 96 well plates containing 1 ml of growth medium prepared by standard procedures. Colonies were picked from the plates and E. coli cultures were grown overnight by standard procedures. Each plate was identified by number. Within each plate, each well contained an individual cDNA clone in the ptrAP vector identified by well position.

Finally, plasmid DNA was extracted from the overnight E. coli cultures using a semi-automated 96-well plasmid DNA miniprep procedure, employing standard procedures for bacterial lysis, genomic DNA precipitation and plasmid DNA purification.

The plasmid DNA extraction was performed as follows:

(a) E. coli were centrifuged for 20 minutes using a Beckman Centrifuge at 3200 rpm.

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(b) Supernatant was discarded and E. coli pellets were resuspended in 130  $\mu$ l WP1 (50 mM TRIS (pH 7.5), 10 mM EDTA, 100  $\mu$ g/ml RNase A) resuspension solution using a TITERTECK MULTIDROP™ apparatus.

5 (c) E. coli pellets were resuspended by vortexing.

(d) 130  $\mu$ l WP2 (0.2 M NaOH, 0.5% SDS) lysing solution was added to each well, and the samples were mixed by vortexing for 5 seconds.

10 (e) 130  $\mu$ l WP3 (125 mM potassium acetate, pH 4.8) neutralizing solution was added to each well, and the samples were mixed by vortexing for 5 seconds.

(f) Samples were placed on ice for 15 minutes, mixed by vortexing for 5 seconds, and recentrifuged for 10 minutes at 3200 rpm in a Beckman Centrifuge.

15 (g) Supernatant (crude DNA extract) was transferred from each well of each 96 well plate into a 96 well filter plate (Polyfiltronics) using a TOMTEC/Quadra 96™ transfer apparatus.

20 (h) 480  $\mu$ l of Wizard™ Midiprep DNA Purification Resin (Promega) was added to each well of each plate containing crude DNA extract using a Titertek Multidrop apparatus and the samples were left for 5 minutes.

25 (i) Each 96 well filter plate was placed on a vacuum housing (Polyfiltronics) and the liquid in each well was removed by suction generated by vacuum created with a Lab Port Vacuum pump.

(j) The Wizard Midiprep DNA Purification Resin in each well (to which plasmid DNA was bound) was washed four times with 600  $\mu$ l of Wizard Wash™.

30 (k) Plates were centrifuged for 5 minutes to remove excessive moisture from the Wizard Midiprep DNA Purification Resin.

(l) Purified plasmid DNAs were eluted from the Wizard Midiprep DNA Purification Resin into collection  
35 plates by addition of 50  $\mu$ l deionized water to each well



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using a Multidrop 8 Channel Pipette, incubation at room temperature for 15 minutes, and centrifugation for 5 minutes (3200 rpm, Beckman centrifuge).

This process resulted in preparation of plasmid DNA contained in 96 well plates with each well containing an individual cDNA clone ligated in the ptrAP expression vector. Individual clones were identified by plate number and well position.

Step 4 Transfection of DNAs into COS7 cells

10 To determine which of the cDNA clones contained within the cDNA library encoded functional signal peptides, individual plasmid DNA preparations were transfected into COS7 cells as follows.

For each 96 well plate of DNA preparations, one 96 well tissue culture plate containing approximately 10,000 COS7 cells per well was prepared using standard procedures.

Immediately prior to DNA transfection, the COS7 cell culture medium in each well of each 96 well plate was replaced with 80  $\mu$ l of OptiMEM (Gibco-BRL; catalog #31985-021) containing 1  $\mu$ l of lipofectamine (Gibco-BRL) and 2  $\mu$ l (approximately 100-200 ng) of DNA prepared as described above. Thus, each well of each 96 well plate containing COS7 cells received DNA representing one individual cDNA clone from the cDNA library in ptrAP3. The COS7 cells were incubated with the Opti-MEM/Lipofectamine/DNA mixture overnight to allow transfection of cells with the plasmid DNAs.

After overnight incubation, the transfection medium was removed from the cells and replaced with 80  $\mu$ l fresh medium composed of Opti-MEM + 1% fetal calf serum. Cells were incubated overnight.

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Step 5 Alkaline Phosphatase Assay

The secreted alkaline phosphatase activity of the transfected COS7 cells was measured as follows. Samples (10  $\mu$ l) of supernatants from the transfected COS7 cells were transferred from each well of each 96 well plate into one well of a Microfluor scintillation plate (Dynatech:Location Catalog #011-010-7805). AP activity in the supernatants was determined using the Phospha-Light Kit (Tropix Inc.; catalog #BP300). AP assays were performed according to the manufacturer's instruction using a Wallace Micro-Beta scintillation counter.

Step 6 Sequencing and Analysis of Positive Clones

The individual plasmid DNAs scoring positive in the COS7 cell AP secretion assay were analyzed further by DNA sequencing using standard procedures. The resulting DNA sequence information was used to perform BLAST sequence similarity searches of nucleotide protein databases to ascertain whether the clone in question encodes either 1) a known secreted or membrane-associated protein possessing a signal sequence, or 2) a putative novel, secreted or membrane-associated protein possessing a putative novel signal sequence.

Identification of the Protein Tyrosine Phosphatase Sigma (PTP $\sigma$ ) Signal Sequence by Mammalian Signal Peptide trAP

Employing the method described in Example 1, a cDNA clone designated ethb005c07 was found to score positive in the COS7 cell transfection AP assay. BLAST similarity searching with the DNA sequence from this clone identified ethb005c07 as a cDNA encoding the signal sequence of protein tyrosine phosphatase sigma (PTP $\sigma$ ), a previously described protein that is well established in the scientific literature to be a transmembrane prot in

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(Pulido et al., Proc. Nat'l Acad. Sci. USA 92:11686, 1995).

Identification of a Novel Immunoglobulin Domain  
Containing Protein by Mammalian Signal Peptide trAP

5           Employing the method described in Example 1, a cDNA clone designated ethb0018f2 was found to score positive in the COS7 cell transfection AP assay. DNA sequencing revealed that ethb0018f2 harbors a 1455 base pair cDNA having a single open reading frame commencing  
10 at nucleotide 55 and continuing to nucleotide 1455. Thus, the ethb0018f2 cDNA encodes a 467 amino acid open reading frame (FIG. 5, SEQ ID NO:5) fused to the AP reporter. Inspection of the ethb0018f2 protein sequence revealed the presence of a putative signal sequence  
15 between amino acids 1 to 20, predicted by the signal peptide prediction algorithm, signal P (Von Heijne, Nucleic Acids. Reg. 14:4683-90, 1986). Thus, ethb0018f2 encodes a partial clone of a novel putative secreted/membrane protein. BLAST similarity searching of  
20 nucleic acid and protein databases with the ethb0018f2 DNA sequence from this clone revealed similarity to a family of proteins known to contain a protein motif referred to as an Immunoglobulin of IgG domain.

          Further visual inspection of the ethb0018f2  
25 protein sequence resulted in the identification of 5 consecutive IgG repeats, defined by a conserved spacing of cysteine, tryptophan, tyrosine, and cysteine residues (FIG. 5).

          FIG. 6 is a depiction of a protein sequence  
30 alignment between clone ethb0018f2 (referred to as 8f2) and seven related proteins known to contain IgG domains that are also known to be expressed in the brain. These proteins are rat neural adhesion molecule f3 (D38492), Drosophila Neuroglial (P20241), human neural adhesion

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molecule L1 (P32004), chick neural adhesion molecule related (P35331), human Axonin 1 (Q02246), rat neural adhesion molecule BIG1 (U11031) and chicken Neurofascin (X65224). Given this sequence similarity, it is likely  
5 that clone ethb0018f2 represents a partial cDNA clone representing a novel protein, expressed in the brain, which contains multiple, consecutive IgG domains. Specifically, since the closest relatives of clone ethb0018f2 are believed to function as neural adhesion  
10 molecules, it is likely that clone ethb0018f2 represents a partial cDNA clone of a novel neural adhesion molecule.

#### Other Embodiments

It is to be understood that while the invention has been described in conjunction with the detailed  
15 description thereof, that the foregoing description is intended to illustrate and not limit the scope of the invention, which is defined by the scope of the appended claims.

## SEQUENCE LISTING

## (1) GENERAL INFORMATION

- (i) APPLICANT: Millennium Biotherapeutics, Inc.
- (ii) TITLE OF THE INVENTION: METHOD FOR IDENTIFYING GENES  
ENCODING NOVEL SECRETED OR MEMBRANE-ASSOCIATED PROTEIN
- (iii) NUMBER OF SEQUENCES: 14
- (iv) CORRESPONDENCE ADDRESS:
  - (A) ADDRESSEE: Fish & Richardson, P.C.
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  - (D) STATE: MA
  - (E) COUNTRY: US
  - (F) ZIP: 02110-2804
- (v) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE: Diskette
  - (B) COMPUTER: IBM Compatible
  - (C) OPERATING SYSTEM: Windows95
  - (D) SOFTWARE: FastSEQ for Windows Version 2.0
- (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER: PCT/US97/-----
  - (B) FILING DATE: 04-NOV-1997
  - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
  - (A) APPLICATION NUMBER: 08/752,307
  - (B) FILING DATE: 19-NOV-1996
- (viii) ATTORNEY/AGENT INFORMATION:
  - (A) NAME: Meiklejohn, Ph.D., Anita L.
  - (B) REGISTRATION NUMBER: 35,283
  - (C) REFERENCE/DOCKET NUMBER: 09404/020W01
- (ix) TELECOMMUNICATION INFORMATION:
  - (A) TELEPHONE: 617-542-5070
  - (B) TELEFAX: 617-542-8906
  - (C) TELEX: 200154

## (2) INFORMATION FOR SEQ ID NO:1:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 4951 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

AAGCTTGGCT	GTGGAATGTG	TGTCAGTTAG	GGTGTGGAAA	GTCCCCAGGC	TCCCCAGCAG	60
GCAGAAGTAT	GCAAAGCATG	CATCTCAATT	AGTCAGCAAC	CAGGTGTGGA	AAGTCCCCAG	120
GCTCCCCAGC	AGGCAGAAGT	ATGCAAAGCA	TGCATCTCAA	TTAGTCAGCA	ACCATAGTCC	180
CGCCCCTAAC	TCCGCCCATC	CCGCCCTAA	CTCCGCCCAG	TTCCGCCCAT	TCTCCGCCCC	240
ATGGCTGACT	AATTTTTTTT	ATTTATGCAG	AGGCCGAGGC	CGCCTCGGCC	TCTGAGCTAT	300
TCCAGAAGTA	GTGAGGAGGC	TTTTTTGGAG	GCCTAGGCTT	TTGCAAAAAG	CTCCTCCGAT	360
CGAGGGGCTC	GCATCTCTCC	TTACGCGGCC	CGCCGCCCTA	CCTGAGGCCG	CCATCCACGC	420
CGGTTGAGTC	GCGTTCTGCC	GCCTCCCGCC	TGTGGTGCCT	CCTGAACTGC	GTCCGCCGTC	480
TAGGTAAGTT	TAAAGCTCAG	GTCGAGACCG	GGCCTTTGTC	CGGCGCTCCC	TTGGAGCCTA	540
CCTAGACTCA	GCCGGCTCTC	CACGCTTTGC	CTGACCCCTG	TTGCTCAACT	CTACGTCTTT	600
GTTTCGTTTT	CTGTTCTGCG	CCGTTACAGA	TCCAAGCTCT	GAAAAACCAG	AAAGTTAACT	660
GGTAAGTTTA	GTCTTTTTGT	CTTTTATTTT	AGGTCCCAGG	TCCCGGATCC	GGTGATCCAA	720
ATCTAAGAAC	TGCTCCTCAG	TGAGTGTTGC	CTTTACTTCT	AGGCCTGTAC	GGAAAGTGTTA	780
CTTCTGCTCT	AAAAGCTGCG	GAATTCGCAC	CACCGTAGTT	TTTACGCCCG	GTGAGCGCTC	840

CACCCGCAACC	TACAAGCGCG	TGTATGATGA	GGTGTACGGC	GACGAGGACC	TGCTTGAGCA	900
GGCCAACGAG	CGCCTCGGGG	AGTTTGCCTA	CGGAAAGCGG	CATAAGGACA	TGTTGGCGTT	960
GCCGCTGGAC	GAGGGCAACC	CAACACCTAG	CCTAAAGCCC	GTGACACTGC	AGCAGGTGCT	1020
GCCCACGCTT	GCACCGTCCG	AAGAAAAGCG	CGGCCATAAG	CGCGAGTCTG	GTGACTTGCC	1080
ACCCACCGTG	CAGCTGATGG	TACCCAAGCG	CCAGCGACTG	GAAGATGTCT	TGGAAAAAAT	1140
GACCGTGGAG	CCTGGGCTGG	AGCCCCAGGT	CCGCGTGCGG	CCAATCAAGC	AGGTGGCACC	1200
GGGACTGGGC	GTGCAGACCG	TGGACGTTCA	GATACCCACC	ACCAGTAGCA	CTAGTATTGC	1260
CACTTGCCACA	GAGGGCATGG	AGACACAAAC	GTCCCCGGTT	GCCTAGCTCG	AGATCATCCC	1320
AGTTGAGGAG	GAGAACCCCG	ACTTCTGGAA	CCGCGAGGCA	GCCGAGGCCC	TGGGTGCCGC	1380
CAAGAAGCTG	CAGCCTGCAC	AGACAGCCGC	CAAGAACCTC	ATCATCTTCC	TGGGCGATGG	1440
GATGGGGGTG	TCTACGGTGA	CAGCTGCCAG	GATCCTAAAA	GGGCAGAAGA	AGGACAAACT	1500
GGGGCCTGAG	ATACCCCTGG	CCATGGACCG	CTTCCCATAT	GTGGCTCTGT	CCAAGACATA	1560
CAATGTAGAC	AAACATGTGC	CAGACAGTGG	AGCCACAGCC	ACGGCCTACC	TGTGCGGGGT	1620
CAAGGGCAAC	TTCCAGAGCA	TTGGCTTGAG	TGCAGCCGCC	CGCTTTAACC	AGTGCAACAC	1680
GACACGCGGC	AACGAGGTCA	TCTCCGTGAT	GAATCGGGCC	AAGAAAGCAG	GGAAGTCAGT	1740
GGGAGTGGTA	ACCACCACAC	GAGTGCAGCA	CGCCTCGCCA	GCCGGCACCT	ACGCCCACAC	1800
GGTGAACCGC	AACTGGTACT	CGGACGCCGA	CGTGCCCTGC	TCGGCCCCGC	AGGAGGGGTG	1860
CCAGGACATC	CTACGCGAGC	TCATCTCCAA	CATGGACATT	GACGTGATCC	TAGGTGGAGG	1920
CCGAAAGTAC	ATGTTTCGCA	TGGGAACCCC	AGACCCTGAG	TACCCAGATG	ACTACAGCCA	1980
AGGTGGGACC	AGGCTGGACG	GGAAGAATCT	GGTGCAGGAA	TGGCTGGCGA	AGCGCCAGGG	2040
TGCCCCGGTAT	GTGTGGAACC	GCACTGAGCT	CATGCAGGCT	TCCCTGGACC	CGTCTGTGAC	2100
CCATCTCTTG	GGTCTCTTTG	AGCCTGGAGA	CATGAAATCA	GAGATCCACC	GAGACTCCAC	2160
ACTGGACCCC	TCCCTGATGG	AGATGACAGA	GGTGCCTCTG	CGCCTGCTGA	CGAGGAACCC	2220
CCGCGGCTTC	TTCCTCTTCG	TGGAGGGTGG	TCGCATCGAC	CATGGTCATC	ATGAAAGCAG	2280
GGCTTACCGG	GCACTGACTG	AGACGATCAT	GTTTCGACGAC	GCCATTGAGA	GGGCGGGCCA	2340
GCTCACCAGC	GAGGAGGACA	CGCTGAGCCT	CGTCACTGCC	GACCACTCCC	ACGTCTTCTC	2400
CTTCGGAGGC	TACCCCTGCG	GAGGGAGCTC	CATCTTCGGG	CTGGCCCCCTG	GCAAGGCCCG	2460
GGACAGGAAG	GCCTACACGG	TCCTCCTATA	CGGAAACGGT	CCAGGCTATG	TGCTCAAGGA	2520
CGGCGCCCGG	CCGGATGTTA	CCGAGAGCGA	GAGCGGGAGC	CCCGAGTATC	GGCAGCAGTC	2580
AGCAGTGCCC	CTGGACGAAG	AGACCCACGC	AGGCGAGGAC	GTGGCGGTGT	TCGCGCGCGG	2640
CCCGCAGGCC	CACCTGGTTC	ACGGCGTGCA	GGAGCAGACC	TTCATAGCGC	ACGTCATGGC	2700
CTTCGCGCGC	TGCGTGAGGC	CCTACACCGC	CTCGCACCTG	GCGCCCCCGG	CTCCGCCACC	2760
CGACGCCGCG	CACCCGGGTT	GAAGTAGTCT	AGAGAAAAAA	CCTCCCACAC	CTCCCCCTGA	2820
ACCTGAAACA	TAAATGAAT	GCAATTGTTG	TTGTTAACTT	GTTTATTGCA	GCTTATAATG	2880
GTTACAAATA	AAGCAATAGC	ATCACAATTT	TCACAAATAA	AGCATTTTTT	TCAGTGCATT	2940
CTAGTTGTGG	TTTGTCCTAA	CTCATCAATG	TATCTTATCA	TGTCTGGATC	CCCGGGTACC	3000
GAGCTCGAAT	TAATTCTCTT	TCCGCTTCCT	CGTCACTGTA	CTCGCTGCGC	TCGGTCTGTC	3060
GGCTGCGGCG	AGCGGTATCA	GCTCACTCAA	AGGCGGTAAT	ACGGTTATCC	ACAGAATCAG	3120
GGGATAACGC	AGGAAAGAAC	ATGTGAGCAA	AAGGCCAGCA	AAAGGCCAGG	AACCGTAAAA	3180
AGGCCGCGTT	GCTGGCGTTC	TTCCATAGCG	TCCGCCCCCC	TGACGAGCAT	CACAAAAATC	3240
GACGCTCAAG	TCAGAGGTGG	CGAAACCCGA	CGAGCATATA	AAGTACCAG	GCGTTTCCCC	3300
CTGGAAGCTC	CCTCGTGCGC	TCTCCTGTTC	CGACCCTGCC	GCTTACCGGA	TACCTGTCCG	3360
CCTTTCTCCC	TTCGGGAAGC	GTGGCGCTTT	CTCAATGCTC	ACGCTGTAGG	TATCTCAGTT	3420
CGGTGTAGGT	CGTTCGCTCC	AAGCTGGGCT	GTGTGCACGA	ACCCCCCGTT	CAGCCCGACC	3480
GCTGCGCCTT	ATCCGGTAAC	TATCGTCTTG	AGTCCAACCC	GGTAAGACAC	GACTTATCGC	3540
CACTGGCAGC	AGCCACTGGT	AACAGGATTA	GCAGAGCGAG	GTATGTAGGC	GGTGTACAG	3600
AGTTCTTGAA	GTGGTGGCCT	AACTACGGCT	ACACTAGAAG	GACAGTATTT	GGTATCTGCG	3660
CTCTGCTGAA	GCCAGTTACC	TTCGGAAAAA	GAGTTGGTAG	CTCTTGATCC	GGCAACAAA	3720
CCACCGCTGG	TAGCGGTGGT	TTTTTTGTTT	GCAAGCAGCA	GATTACGCGC	AGAAAAAAG	3780
GATCTCAAGA	AGATCCTTTG	ATCTTTCTTA	CGGGTCTGTA	CGCTCAGTGG	AACGAAAAC	3840
CACGTTAAGG	GATTTTGGTC	ATGAGATTAT	CAAAAAGGAT	CTTCACCTAG	ATCCTTTTAA	3900
ATTAAAAATG	AAGTTTAAAA	TCAATCTAAA	GTATATATGA	GTAAACTTGG	TCTGACAGTT	3960
ACCAATGCTT	AATCAGTGAG	GCACCTATCT	CAGCGATCTG	TCTATTTCTG	TCATCCATAG	4020
TTGCCGTGACT	CCCCGTCGTG	TAGATAACTA	CGATACGGGA	GGGCTTACCA	TCTGGCCCCA	4080
GTGCTGCAAT	GATACCGCGA	GACCCACGCT	CACCGGCTCC	AGATTTATCA	GCAATAAACC	4140
AGCCAGCCCG	AAGGGCCGAG	CGCAGAAGTG	GTCCTGCAAC	TTTATCCGCC	TCCATCCAGT	4200
CTATTAATTG	TTGCCGGGAA	GCTAGAGTAA	GTAGTTCGCC	AGTTAATAGT	TTGCGCAACG	4260
TTGTTGCCAT	TGCTACAGGC	ATCGTGGTGT	CACGCTCGTC	GTTTGGTATG	GCTTCATTCA	4320
GCTCCGGTTC	CCAACGATTA	AGGCGAGTTA	CATGATCCCC	CATGTTGTGC	AAAAAGCCGG	4380
TTAGCTCCTT	CGGTCCCTCCG	ATCGTTGTCA	GAAGTAAGTT	GGCCGCGAGT	TTATCACTCA	4440
TGTTTATGGC	AGCACTGCAT	AATTCTCTTA	CTGTCTATGC	ATCCGTAAGA	TGCTTTTCTG	4500
TGACTGGTGA	GTAATCAACC	AAGTCATTCT	GAGAATAGTG	TATGCGGCGA	CCGAGTTGCT	4560
CTTGCCCGGC	GTCAATACGG	GATAATACCG	CGCCACATAG	CAGAACTTTA	AAAGTGCTCA	4620
TCATTGGAAA	ACGTTCTTCG	GGGCGAAAAA	TCTCAAGGAT	CTTACCGCTG	TTGAGATCCA	4680
GTTTCGATGTA	ACCCACTCGT	GCACCCAACT	GATCTTCAGC	ATCTTTTACT	TTCACCGACG	4740
TTTCTGGGTG	AGCAAAAACA	GGAAGGCAAA	ATGCCGCAAA	AAAGGGAATA	AGGGCGACAC	4800
GGAAATGTTG	AATACTCATA	CTCTTCCTTT	TTCAATATTA	TTGAAGCATT	TATCAGGGTT	4860
ATTGTCTCAT	GAGCGGATAC	ATATTTGAAT	GTATTTAGAA	AAATAAACAA	ATAGGGGTTT	4920
CGCGCACATT	TCCCCGAAAA	GTGCCACCTG	C			4951

## (2) INFORMATION FOR SEQ ID NO:2:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 530 amino acids  
 (B) TYPE: amino acid  
 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

```

Met Leu Leu Leu Leu Leu Leu Gly Leu Arg Leu Gln Leu Ser Leu
 1      5      10      15
Gly Ile Ile Pro Val Glu Glu Glu Asn Pro Asp Phe Trp Asn Arg Glu
 20      25      30
Ala Ala Glu Ala Leu Gly Ala Ala Lys Lys Leu Gln Pro Ala Gln Thr
 35      40      45
Ala Ala Lys Asn Leu Ile Ile Phe Leu Gly Asp Gly Met Gly Val Ser
 50      55      60
Thr Val Thr Ala Ala Arg Ile Leu Lys Gly Gln Lys Lys Asp Lys Leu
 65      70      75      80
Gly Pro Glu Ile Pro Leu Ala Met Asp Arg Phe Pro Tyr Val Ala Leu
 85      90      95
Ser Lys Thr Tyr Asn Val Asp Lys His Val Pro Asp Ser Gly Ala Thr
100      105      110
Ala Thr Ala Tyr Leu Cys Gly Val Lys Gly Asn Phe Gln Thr Ile Gly
115      120      125
Leu Ser Ala Ala Ala Arg Phe Asn Gln Cys Asn Thr Thr Arg Gly Asn
130      135      140
Glu Val Ile Ser Val Met Asn Arg Ala Lys Lys Ala Gly Lys Ser Val
145      150      155      160
Gly Val Val Thr Thr Thr Arg Val Gln His Ala Ser Pro Ala Gly Thr
165      170      175
Tyr Ala His Thr Val Asn Arg Asn Trp Tyr Ser Asp Ala Asp Val Pro
180      185      190
Ala Ser Ala Arg Gln Glu Gly Cys Gln Asp Ile Ala Thr Gln Leu Ile
195      200      205
Ser Asn Met Asp Ile Asp Val Ile Leu Gly Gly Gly Arg Lys Tyr Met
210      215      220
Phe Arg Met Gly Thr Pro Asp Pro Glu Tyr Pro Asp Asp Tyr Ser Gln
225      230      235      240
Gly Gly Thr Arg Leu Asp Gly Lys Asn Leu Val Gln Glu Trp Leu Ala
245      250      255
Lys Arg Gln Gly Ala Arg Tyr Val Trp Asn Arg Thr Glu Leu Met Gln
260      265      270
Ala Ser Leu Asp Pro Ser Val Thr His Leu Met Gly Leu Phe Glu Pro
275      280      285
Gly Asp Met Lys Tyr Glu Ile His Arg Asp Ser Thr Leu Asp Pro Ser
290      295      300
Leu Met Glu Met Thr Glu Ala Ala Leu Arg Leu Leu Ser Arg Asn Pro
305      310      315      320
Arg Gly Phe Phe Leu Phe Val Glu Gly Gly Arg Ile Asp His Gly His
325      330      335
His Glu Ser Arg Ala Tyr Arg Ala Leu Thr Glu Thr Ile Met Phe Asp
340      345      350
Asp Ala Ile Glu Arg Ala Gly Gln Leu Thr Ser Glu Glu Asp Thr Leu
355      360      365
Ser Leu Val Thr Ala Asp His Ser His Val Phe Ser Phe Gly Gly Tyr
370      375      380
Pro Leu Arg Gly Ser Ser Ile Phe Gly Leu Ala Pro Gly Lys Ala Arg
385      390      395      400
Asp Arg Lys Ala Tyr Thr Val Leu Leu Tyr Gly Asn Gly Pro Gly Tyr
405      410      415
Val Leu Lys Asp Gly Ala Arg Pro Asp Val Thr Glu Ser Glu Ser Gly
420      425      430
Ser Pro Glu Tyr Arg Gln Gln Ser Ala Val Pro Leu Asp Glu Thr
435      440      445
His Ala Gly Glu Asp Val Ala Val Phe Ala Arg Gly Pro Gln Ala His
450      455      460

```

Leu Val His Gly Val Gln Glu Gln Thr Phe Ile Ala His Val Met Ala  
 465 470 475 480  
 Phe Ala Ala Cys Leu Glu Pro Tyr Thr Ala Cys Asp Leu Ala Pro Pro  
 485 490 495  
 Ala Gly Thr Thr Asp Ala Ala His Pro Gly Arg Ser Val Val Pro Ala  
 500 505 510  
 Leu Leu Pro Leu Leu Ala Gly Thr Leu Leu Leu Leu Glu Thr Ala Thr  
 515 520 525  
 Ala Pro  
 530

## (2) INFORMATION FOR SEQ ID NO:3:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 489 amino acids  
 (B) TYPE: amino acid  
 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Ile Ile Pro Val Glu Glu Glu Asn Pro Asp Phe Trp Asn Arg Glu Ala  
 1 5 10 15  
 Ala Glu Ala Leu Gly Ala Ala Lys Lys Leu Gln Pro Ala Gln Thr Ala  
 20 25 30  
 Ala Lys Asn Leu Ile Ile Phe Leu Gly Asp Gly Met Gly Val Ser Thr  
 35 40 45  
 Val Thr Ala Ala Arg Ile Leu Lys Gly Gln Lys Lys Asp Lys Leu Gly  
 50 55 60  
 Pro Glu Ile Pro Leu Ala Met Asp Arg Phe Pro Tyr Val Ala Leu Ser  
 65 70 75 80  
 Lys Thr Tyr Asn Val Asp Lys His Val Pro Asp Ser Gly Ala Thr Ala  
 85 90 95  
 Thr Ala Tyr Leu Cys Gly Val Lys Gly Asn Phe Gln Thr Ile Gly Leu  
 100 105 110  
 Ser Ala Ala Ala Arg Phe Asn Gln Cys Asn Thr Thr Arg Gly Asn Glu  
 115 120 125  
 Val Ile Ser Val Met Asn Arg Ala Lys Lys Ala Gly Lys Ser Val Gly  
 130 135 140  
 Val Val Thr Thr Thr Arg Val Gln His Ala Ser Pro Ala Gly Thr Tyr  
 145 150 155 160  
 Ala His Thr Val Asn Arg Asn Trp Tyr Ser Asp Ala Asp Val Pro Ala  
 165 170 175  
 Ser Ala Arg Gln Glu Gly Cys Gln Asp Ile Ala Thr Gln Leu Ile Ser  
 180 185 190  
 Asn Met Asp Ile Asp Val Ile Leu Gly Gly Gly Arg Lys Tyr Met Phe  
 195 200 205  
 Arg Met Gly Thr Pro Asp Pro Glu Tyr Pro Asp Asp Tyr Ser Gln Gly  
 210 215 220  
 Gly Thr Arg Leu Asp Gly Lys Asn Leu Val Gln Glu Trp Leu Ala Lys  
 225 230 235 240  
 Arg Gln Gly Ala Arg Tyr Val Trp Asn Arg Thr Glu Leu Met Gln Ala  
 245 250 255  
 Ser Leu Asp Pro Ser Val Thr His Leu Met Gly Leu Phe Glu Pro Gly  
 260 265 270  
 Asp Met Lys Tyr Glu Ile His Arg Asp Ser Thr Leu Asp Pro Ser Leu  
 275 280 285  
 Met Glu Met Thr Glu Ala Ala Leu Arg Leu Leu Ser Arg Asn Pro Arg  
 290 295 300  
 Gly Phe Phe Leu Phe Val Glu Gly Gly Arg Ile Asp His Gly His His  
 305 310 315 320  
 Glu Ser Arg Ala Tyr Arg Ala Leu Thr Glu Thr Ile Met Phe Asp Asp  
 325 330 335  
 Ala Ile Glu Arg Ala Gly Gln Leu Thr Ser Glu Glu Asp Thr Leu Ser  
 340 345 350  
 Leu Val Thr Ala Asp His Ser His Val Phe Ser Phe Gly Tyr Pro  
 355 360 365



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```

Leu Arg Gly Ser Ser Ile Phe Gly Leu Ala Pro Gly Lys Ala Arg Asp
 370      375      380
Arg Lys Ala Tyr Thr Val Leu Leu Tyr Gly Asn Gly Pro Gly Tyr Val
 385      390      395      400
Leu Lys Asp Gly Ala Arg Pro Asp Val Thr Glu Ser Glu Ser Gly Ser
      405      410      415
Pro Glu Tyr Arg Gln Gln Ser Ala Val Pro Leu Asp Glu Glu Thr His
      420      425      430
Ala Gly Glu Asp Val Ala Val Phe Ala Arg Gly Pro Gln Ala His Leu
      435      440      445
Val His Gly Val Gln Glu Gln Thr Phe Ile Ala His Val Met Ala Phe
      450      455      460
Ala Ala Cys Leu Glu Pro Tyr Thr Ala Cys Asp Leu Ala Pro Pro Ala
 465      470      475      480
Gly Thr Thr Asp Ala Ala His Pro Gly
      485

```

## (2) INFORMATION FOR SEQ ID NO:4:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 17 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

CTGGACTCGA GNNNNNN

17

## (2) INFORMATION FOR SEQ ID NO:5:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 465 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (v) FRAGMENT TYPE: internal

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

```

Met Trp Leu Val Thr Phe Leu Leu Leu Leu Asp Ser Leu His Lys Ala
 1      5      10      15
Arg Pro Glu Asp Val Gly Thr Ser Leu Tyr Phe Val Asn Asp Ser Leu
      20      25      30
Gln Gln Val Thr Phe Ser Ser Ser Val Gly Val Val Val Pro Cys Pro
      35      40      45
Ala Ala Gly Ser Pro Ser Ala Ala Leu Arg Trp Tyr Leu Ala Thr Gly
      50      55      60
Asp Asp Ile Tyr Asp Val Pro His Ile Arg His Val His Ala Asn Gly
 65      70      75      80
Thr Leu Gln Leu Tyr Pro Phe Ser Pro Ser Ala Phe Asn Ser Phe Ile
      85      90      95
His Asp Asn Asp Tyr Phe Cys Thr Ala Glu Asn Ala Ala Gly Lys Ile
      100      105      110
Arg Ser Pro Asn Ile Arg Val Lys Ala Val Phe Arg Glu Pro Tyr Thr
      115      120      125
Val Arg Val Glu Asp Gln Arg Ser Met Arg Gly Asn Val Ala Val Phe
      130      135      140
Lys Cys Leu Ile Pro Ser Ser Val Gln Glu Tyr Val Ser Val Val Ser
 145      150      155      160
Trp Glu Lys Asp Thr Val Ser Ile Ile Pro Glu Asn Arg Phe Phe Ile
      165      170      175
Thr Tyr His Gly Leu Tyr Ile Ser Asp Val Gln Lys Glu Asp Ala
      180      185      190

```

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```

Leu Ser Thr Tyr Arg Cys Ile Thr Lys His Lys Tyr Ser Gly Glu Thr
    195                200                205
Arg Gln Ser Asn Gly Ala Arg Leu Ser Val Thr Asp Pro Ala Glu Ser
    210                215                220
Ile Pro Thr Ile Leu Asp Gly Phe His Ser Gln Glu Val Trp Ala Gly
225                230                235                240
His Thr Val Glu Leu Pro Cys Thr Ala Ser Gly Tyr Pro Ile Pro Ala
    245                250                255
Ile Arg Trp Leu Lys Asp Gly Arg Pro Leu Pro Ala Asp Ser Arg Trp
    260                265                270
Thr Lys Arg Ile Thr Gly Leu Thr Ile Ser Asp Leu Arg Thr Glu Asp
    275                280                285
Ser Gly Thr Tyr Ile Cys Glu Val Thr Asn Thr Phe Gly Ser Ala Glu
    290                295                300
Ala Thr Gly Ile Leu Met Val Ile Asp Pro Leu His Val Thr Leu Thr
305                310                315                320
Pro Lys Lys Leu Lys Thr Gly Ile Gly Ser Thr Val Ile Leu Ser Cys
    325                330                335
Ala Leu Thr Gly Ser Pro Glu Phe Thr Ile Arg Trp Tyr Arg Asn Thr
    340                345                350
Glu Leu Val Leu Pro Asp Glu Ala Ile Ser Ile Arg Gly Leu Ser Asn
    355                360                365
Glu Thr Leu Leu Ile Thr Ser Ala Gln Lys Ser His Ser Gly Ala Tyr
    370                375                380
Gln Cys Phe Ala Thr Arg Lys Ala Gln Thr Ala Gln Asp Phe Ala Ile
385                390                395                400
Ile Ala Leu Glu Asp Gly Thr Pro Arg Ile Val Ser Ser Phe Ser Glu
    405                410                415
Lys Val Val Asn Pro Gly Glu Gln Phe Ser Leu Met Cys Ala Ala Lys
    420                425                430
Gly Ala Pro Pro Pro Thr Val Thr Trp Ala Leu Asp Asp Glu Pro Ile
    435                440                445
Val Arg Asp Gly Ser His Arg Thr Asn Gln Tyr Thr Met Ser Asp Gly
    450                455                460
Thr
465

```

## (2) INFORMATION FOR SEQ ID NO:6:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 1493 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: cDNA

## (ix) FEATURE:

- (A) NAME/KEY: Coding Sequence
- (B) LOCATION: 99...1493

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

```

GGCACGAGGG CGGCTGGGAG CGCGCTGAGC GGGGGAGAGG CGCTGCCGCA CGGCCGGCCA      60
CAGGACCACC TCCCCGAGA ATAGGGCCTC TTTATGGC ATG TGG CTG GTA ACT TTC      116
                                   Met Trp Leu Val Thr Phe
                                   1               5

CTC CTG CTC CTG GAC TCT TTA CAC AAA GCC CGC CCT GAA GAT GTT GGC      164
Leu Leu Leu Leu Asp Ser Leu His Lys Ala Arg Pro Glu Asp Val Gly
    10                15                20

ACC AGC CTC TAC TTT GTA AAT GAC TCC TTG CAG CAG GTG ACC TTT TCC      212
Thr Ser Leu Tyr Phe Val Asn Asp Ser Leu Gln Gln Val Thr Phe Ser
    25                30                35

AGC TCC GTG GGG GTG GTG GTG CCC TGC CCG GCC GCG GGC TCC CCC AGC      260
Ser Ser Val Gly Val Val Val Pro Cys Pro Ala Ala Gly Ser Pro Ser
    40                45                50

```

GCG	GCC	CTT	CGA	TGG	TAC	CTG	GCC	ACA	GGG	GAC	GAC	ATC	TAC	GAC	GTG	308
Ala	Ala	Leu	Arg	Trp	Tyr	Leu	Ala	Thr	Gly	Asp	Asp	Ile	Tyr	Asp	Val	
55					60					65					70	
CCG	CAC	ATC	CGG	CAC	GTC	CAC	GCC	AAC	GGG	ACG	CTG	CAG	CTC	TAC	CCC	356
Pro	His	Ile	Arg	His	Val	His	Ala	Asn	Gly	Thr	Leu	Gln	Leu	Tyr	Pro	
				75					80					85		
TTC	TCC	CCC	TCC	GCC	TTC	AAT	AGC	TTT	ATC	CAC	GAC	AAT	GAC	TAC	TTC	404
Phe	Ser	Pro	Ser	Ala	Phe	Asn	Ser	Phe	Ile	His	Asp	Asn	Asp	Tyr	Phe	
			90					95					100			
TGC	ACC	GCG	GAG	AAC	GCT	GCC	GGC	AAG	ATC	CGG	AGC	CCC	AAC	ATC	CGC	452
Cys	Thr	Ala	Glu	Asn	Ala	Ala	Gly	Lys	Ile	Arg	Ser	Pro	Asn	Ile	Arg	
		105					110					115				
GTC	AAA	GCA	GTT	TTC	AGG	GAA	CCC	TAC	ACC	GTC	CGG	GTG	GAG	GAT	CAA	500
Val	Lys	Ala	Val	Phe	Arg	Glu	Pro	Tyr	Thr	Val	Arg	Val	Glu	Asp	Gln	
	120					125					130					
AGG	TCA	ATG	CGT	GGC	AAC	GTG	GCC	GTC	TTC	AAG	TGC	CTC	ATC	CCC	TCT	548
Arg	Ser	Met	Arg	Gly	Asn	Val	Ala	Val	Phe	Lys	Cys	Leu	Ile	Pro	Ser	
					140					145					150	
TCA	GTG	CAG	GAA	TAT	GTT	AGC	GTT	GTA	TCT	TGG	GAG	AAA	GAC	ACA	GTC	596
Ser	Val	Gln	Glu	Tyr	Val	Ser	Val	Val	Ser	Trp	Glu	Lys	Asp	Thr	Val	
				155					160					165		
TCC	ATC	ATC	CCA	GAA	AAC	AGG	TTT	TTT	ATT	ACC	TAC	CAC	GGC	GGG	CTG	644
Ser	Ile	Ile	Pro	Glu	Asn	Arg	Phe	Phe	Ile	Thr	Tyr	His	Gly	Gly	Leu	
			170					175					180			
TAC	ATC	TCT	GAC	GTA	CAG	AAG	GAG	GAC	GCC	CTC	TCC	ACC	TAT	CGC	TGC	692
Tyr	Ile	Ser	Asp	Val	Gln	Lys	Glu	Asp	Ala	Leu	Ser	Thr	Tyr	Arg	Cys	
		185					190					195				
ATC	ACC	AAG	CAC	AAG	TAT	AGC	GGG	GAG	ACC	CGG	CAG	AGC	AAT	GGG	GCA	740
Ile	Thr	Lys	His	Lys	Tyr	Ser	Gly	Glu	Thr	Arg	Gln	Ser	Asn	Gly	Ala	
	200					205					210					
CGC	CTC	TCT	GTG	ACA	GAC	CCT	GCT	GAG	TCG	ATC	CCC	ACC	ATC	CTG	GAT	788
Arg	Leu	Ser	Val	Thr	Asp	Pro	Ala	Glu	Ser	Ile	Pro	Thr	Ile	Leu	Asp	
	215				220					225				230		
GGC	TTC	CAC	TCC	CAG	GAA	GTG	TGG	GCC	GGC	CAC	ACC	GTG	GAG	CTG	CCC	836
Gly	Phe	His	Ser	Gln	Glu	Val	Trp	Ala	Gly	His	Thr	Val	Glu	Leu	Pro	
				235					240					245		
TGC	ACC	GCC	TCG	GGC	TAC	CCT	ATC	CCC	GCC	ATC	CGC	TGG	CTC	AAG	GAT	884
Cys	Thr	Ala	Ser	Gly	Tyr	Pro	Ile	Pro	Ala	Ile	Arg	Trp	Leu	Lys	Asp	
			250					255					260			
GGC	CGG	CCC	CTC	CCG	GCT	GAC	AGC	CGC	TGG	ACC	AAG	CGC	ATC	ACA	GGG	932
Gly	Arg	Pro	Leu	Pro	Ala	Asp	Ser	Arg	Trp	Thr	Lys	Arg	Ile	Thr	Gly	
		265					270					275				
CTG	ACC	ATC	AGC	GAC	TTG	CGG	ACC	GAG	GAC	AGC	GGC	ACC	TAC	ATT	TGT	980
Leu	Thr	Ile	Ser	Asp	Leu	Arg	Thr	Glu	Asp	Ser	Gly	Thr	Tyr	Ile	Cys	
	280					285					290					
GAG	GTC	ACC	AAC	ACC	TTC	GGT	TCG	GCA	GAG	GCC	ACA	GGC	ATC	CTC	ATG	1028
Glu	Val	Thr	Asn	Thr	Phe	Gly	Ser	Ala	Glu	Ala	Thr	Gly	Ile	Leu	Met	
	295				300					305					310	
GTC	ATT	GAT	CCC	CTT	CAT	GTG	ACC	CTG	ACA	CCA	AAG	AAG	CTG	AAG	ACC	1076
Val	Ile	Asp	Pro	Leu	His	Val	Thr	Leu	Thr	Pro	Lys	Lys	Leu	Lys	Thr	
				315					320					325		

GGC ATT GGC AGC ACG GTC ATC CTC TCC TGT GCC CTG ACG GGC TCC CCA	1124
Gly Ile Gly Ser Thr Val Ile Leu Ser Cys Ala Leu Thr Gly Ser Pro	
330 335 340	
GAG TTC ACC ATC CGC TGG TAT CGC AAC ACG GAG CTG GTG CTG CCT GAC	1172
Glu Phe Thr Ile Arg Trp Tyr Arg Asn Thr Glu Leu Val Leu Pro Asp	
345 350 355	
GAG GCC ATC TCC ATC CGT GGG CTC AGC AAC GAG ACG CTG CTC ATC ACC	1220
Glu Ala Ile Ser Ile Arg Gly Leu Ser Asn Glu Thr Leu Leu Ile Thr	
360 365 370	
TCG GCC CAG AAG AGC CAT TCC GGG GCC TAC CAG TGC TTC GCT ACC CGC	1268
Ser Ala Gln Lys Ser His Ser Gly Ala Tyr Gln Cys Phe Ala Thr Arg	
375 380 385 390	
AAG GCC CAG ACC GCC CAG GAC TTT GCC ATC ATT GCA CTT GAG GAT GGC	1316
Lys Ala Gln Thr Ala Gln Asp Phe Ala Ile Ala Leu Glu Asp Gly	
395 400 405	
ACG CCC CGC ATC GTC TCG TCC TTC AGC GAG AAG GTG GTC AAC CCC GGG	1364
Thr Pro Arg Ile Val Ser Ser Phe Ser Glu Lys Val Val Asn Pro Gly	
410 415 420	
GAG CAG TTC TCA CTG ATG TGT GCG GCC AAG GGC GCC CCG CCC CCC ACG	1412
Glu Gln Phe Ser Leu Met Cys Ala Ala Lys Gly Ala Pro Pro Pro Thr	
425 430 435	
GTC ACC TGG GCC CTC GAC GAT GAG CCC ATC GTG CGG GAT GGC AGC CAC	1460
Val Thr Trp Ala Leu Asp Asp Glu Pro Ile Val Arg Asp Gly Ser His	
440 445 450	
CGC ACC AAC CAG TAC ACC ATG TCG GAC GGC ACC	1493
Arg Thr Asn Gln Tyr Thr Met Ser Asp Gly Thr	
455 460 465	

## (2) INFORMATION FOR SEQ ID NO:7:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 462 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Met Trp Leu Val Thr Phe Leu Leu Leu Leu Asp Ser Leu His Lys Ala	
1 5 10 15	
Arg Pro Glu Asp Val Gly Thr Ser Leu Tyr Phe Val Asn Asp Ser Leu	
20 25 30	
Gln Gln Val Thr Phe Ser Ser Ser Val Gly Val Val Val Pro Cys Pro	
35 40 45	
Ala Ala Gly Ser Pro Ser Ala Ala Leu Arg Trp Tyr Leu Ala Thr Gly	
50 55 60	
Asp Asp Ile Tyr Asp Val Pro His Ile Arg His Val His Ala Asn Gly	
65 70 75 80	
Thr Leu Gln Leu Tyr Pro Phe Ser Pro Ser Ala Phe Asn Ser Phe Ile	
85 90 95	
His Asp Asn Asp Tyr Phe Cys Thr Ala Glu Asn Ala Ala Gly Lys Ile	
100 105 110	
Arg Ser Pro Asn Ile Arg Val Lys Ala Val Phe Arg Glu Pro Tyr Thr	
115 120 125	
Val Arg Val Glu Asp Gln Arg Ser Met Arg Gly Asn Val Ala Val Phe	
130 135 140	
Lys Cys Leu Ile Pro Ser Ser Val Gln Glu Tyr Val Ser Val Val Ser	
145 150 155 160	
Trp Glu Lys Asp Thr Val Ser Ile Ile Pro Glu Asn Arg Phe Phe Ile	
165 170 175	

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Thr	Tyr	His	Gly	Gly	Leu	Tyr	Ile	Ser	Asp	Val	Gln	Lys	Glu	Asp	Ala
			180					185					190		
Leu	Ser	Thr	Tyr	Arg	Cys	Ile	Thr	Lys	His	Lys	Tyr	Ser	Gly	Glu	Thr
		195					200					205			
Arg	Gln	Ser	Asn	Gly	Ala	Arg	Leu	Ser	Val	Thr	Asp	Pro	Ala	Glu	Ser
	210					215					220				
Ile	Pro	Thr	Ile	Leu	Asp	Gly	Phe	His	Ser	Gln	Glu	Val	Trp	Ala	Gly
225					230					235				240	
His	Thr	Val	Glu	Leu	Pro	Cys	Thr	Ala	Ser	Gly	Tyr	Pro	Ile	Pro	Ala
			245						250					255	
Ile	Arg	Trp	Leu	Lys	Asp	Gly	Arg	Pro	Leu	Pro	Ala	Asp	Ser	Arg	Trp
		260						265					270		
Thr	Lys	Arg	Ile	Thr	Gly	Leu	Thr	Ile	Ser	Asp	Leu	Arg	Thr	Glu	Asp
		275					280					285			
Ser	Gly	Thr	Tyr	Ile	Cys	Glu	Val	Thr	Asn	Thr	Phe	Gly	Ser	Ala	Glu
	290				295						300				
Ala	Thr	Gly	Ile	Leu	Met	Val	Ile	Asp	Pro	Leu	His	Val	Thr	Leu	Thr
305					310					315				320	
Pro	Lys	Lys	Leu	Lys	Thr	Gly	Ile	Gly	Ser	Thr	Val	Ile	Leu	Ser	Cys
			325						330					335	
Ala	Leu	Thr	Gly	Ser	Pro	Glu	Phe	Thr	Ile	Arg	Trp	Tyr	Arg	Asn	Thr
		340						345					350		
Glu	Leu	Val	Leu	Pro	Asp	Glu	Ala	Ile	Ser	Ile	Arg	Gly	Leu	Ser	Asn
	355						360					365			
Glu	Thr	Leu	Leu	Ile	Thr	Ser	Ala	Gln	Lys	Ser	His	Ser	Gly	Ala	Tyr
	370					375					380				
Gln	Cys	Phe	Ala	Thr	Arg	Lys	Ala	Gln	Thr	Ala	Gln	Asp	Phe	Ala	Ile
385					390					395				400	
Ile	Ala	Leu	Glu	Asp	Gly	Thr	Pro	Arg	Ile	Val	Ser	Ser	Phe	Ser	Glu
			405						410					415	
Lys	Val	Val	Asn	Pro	Gly	Glu	Gln	Phe	Ser	Leu	Met	Cys	Ala	Ala	Lys
			420					425					430		
Gly	Ala	Pro	Pro	Pro	Thr	Val	Thr	Trp	Ala	Leu	Asp	Asp	Glu	Pro	Ile
		435					440					445			
Val	Arg	Asp	Gly	Ser	His	Arg	Thr	Asn	Gln	Tyr	Thr	Met	Ser		
	450					455					460				

## (2) INFORMATION FOR SEQ ID NO:8:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 605 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Met	Lys	Thr	Pro	Leu	Leu	Val	Ser	His	Leu	Leu	Ile	Ser	Leu	Thr
1				5					10				15	
Ser	Cys	Leu	Gly	Glu	Phe	Thr	Trp	His	Arg	Arg	Tyr	Gly	His	Gly
		20						25				30		Val
Ser	Glu	Glu	Asp	Lys	Gly	Phe	Gly	Pro	Ile	Phe	Glu	Glu	Gln	Pro
		35					40					45		Ile
Asn	Thr	Ile	Tyr	Pro	Glu	Glu	Ser	Leu	Glu	Gly	Lys	Val	Ser	Leu
	50					55					60			Asn
Cys	Arg	Ala	Arg	Ala	Ser	Pro	Phe	Pro	Val	Tyr	Lys	Trp	Arg	Met
65				70					75					80
Asn	Gly	Asp	Val	Asp	Leu	Thr	Asn	Asp	Arg	Tyr	Ser	Met	Val	Gly
			85						90				95	Gly
Asn	Leu	Val	Ile	Asn	Asn	Pro	Asp	Lys	Gln	Lys	Asp	Ala	Gly	Ile
		100						105					110	Tyr
Tyr	Cys	L u	Ala	Ser	Asn	Asn	Tyr	Gly	Met	Val	Arg	Ser	Thr	Glu
		115					120					125		Ala
Thr	Leu	Ser	Phe	Gly	Tyr	Leu	Asp	Pro	Phe	Pro	Pro	Glu	Asp	Arg
	130					135						140		Pro

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Glu	Val	Lys	Val	Lys	Glu	Gly	Lys	Gly	Met	Val	Leu	Leu	Cys	Asp	Pro
145					150					155					160
Pro	Tyr	His	Phe	Pro	Asp	Asp	Leu	Ser	Tyr	Arg	Trp	Leu	Leu	Asn	Glu
				165					170						175
Phe	Pro	Val	Phe	Ile	Thr	Met	Asp	Lys	Arg	Arg	Phe	Val	Ser	Gln	Thr
			180					185						190	
Asn	Gly	Asn	Leu	Tyr	Ile	Ala	Asn	Val	Glu	Ser	Ser	Asp	Arg	Gly	Asn
		195					200					205			
Tyr	Ser	Cys	Phe	Val	Ser	Ser	Pro	Ser	Ile	Thr	Lys	Ser	Val	Phe	Ser
	210					215					220				
Lys	Phe	Ile	Pro	Leu	Ile	Pro	Ile	Pro	Glu	Arg	Thr	Thr	Lys	Pro	Tyr
225					230					235					240
Pro	Ala	Asp	Ile	Val	Val	Gln	Phe	Lys	Asp	Ile	Tyr	Thr	Met	Met	Gly
			245						250					255	
Gln	Asn	Val	Thr	Leu	Glu	Cys	Phe	Ala	Leu	Gly	Asn	Pro	Val	Pro	Asp
			260					265					270		
Ile	Arg	Trp	Arg	Lys	Val	Leu	Glu	Pro	Met	Pro	Thr	Thr	Ala	Glu	Ile
	275						280					285			
Ser	Thr	Ser	Gly	Ala	Val	Leu	Lys	Ile	Phe	Asn	Ile	Gln	Leu	Glu	Asp
	290					295					300				
Glu	Gly	Leu	Tyr	Glu	Cys	Glu	Ala	Glu	Asn	Ile	Arg	Gly	Lys	Asp	Lys
305					310					315					320
His	Gln	Ala	Arg	Ile	Tyr	Val	Gln	Ala	Phe	Pro	Glu	Trp	Val	Glu	His
			325						330					335	
Ile	Asn	Asp	Thr	Glu	Val	Asp	Ile	Gly	Ser	Asp	Leu	Tyr	Trp	Pro	Cys
			340					345					350		
Val	Ala	Thr	Gly	Lys	Pro	Ile	Pro	Thr	Ile	Arg	Trp	Leu	Lys	Asn	Gly
		355					360					365			
Tyr	Ala	Tyr	His	Lys	Gly	Glu	Leu	Arg	Leu	Tyr	Asp	Val	Thr	Phe	Glu
	370				375						380				
Asn	Ala	Gly	Met	Tyr	Gln	Cys	Ile	Ala	Glu	Asn	Ala	Tyr	Gly	Thr	Ile
385					390					395					400
Tyr	Ala	Asn	Ala	Glu	Leu	Lys	Ile	Leu	Ala	Leu	Ala	Pro	Thr	Phe	Glu
			405						410					415	
Met	Asn	Pro	Met	Lys	Lys	Lys	Ile	Leu	Ala	Ala	Lys	Gly	Gly	Arg	Val
			420					425					430		
Ile	Ile	Glu	Cys	Lys	Pro	Lys	Ala	Ala	Pro	Lys	Pro	Lys	Phe	Ser	Trp
		435					440					445			
Ser	Lys	Gly	Thr	Glu	Trp	Leu	Val	Asn	Ser	Ser	Arg	Ile	Leu	Ile	Trp
	450					455					460				
Glu	Asp	Gly	Ser	Leu	Glu	Ile	Asn	Asn	Ile	Thr	Arg	Asn	Asp	Gly	Gly
465					470					475					480
Ile	Tyr	Thr	Cys	Phe	Ala	Glu	Asn	Asn	Arg	Gly	Lys	Ala	Asn	Ser	Thr
			485						490					495	
Gly	Thr	Leu	Val	Ile	Thr	Asn	Pro	Thr	Arg	Ile	Ile	Leu	Ala	Pro	Ile
			500					505					510		
Asn	Ala	Asp	Ile	Thr	Val	Gly	Glu	Asn	Ala	Thr	Met	Gln	Cys	Ala	Ala
		515					520					525			
Ser	Phe	Asp	Pro	Ser	Leu	Asp	Leu	Thr	Phe	Val	Trp	Ser	Phe	Asn	Gly
	530					535					540				
Tyr	Val	Ile	Asp	Phe	Asn	Lys	Glu	Ile	Thr	Asn	Ile	His	Tyr	Gln	Arg
545					550					555					560
Asn	Phe	Met	Leu	Asp	Ala	Asn	Gly	Glu	Leu	Ile	Arg	Asn	Ala	Gln	
			565						570					575	
Leu	Lys	His	Ala	Gly	Arg	Tyr	Thr	Cys	Thr	Ala	Gln	Thr	Ile	Val	Asp
			580					585					590		
Asn	Ser	Ser	Ala	Ser	Ala	Asp	Leu	Val	Val	Arg	Gly	Pro			
		595					600					605			

## (2) INFORMATION FOR SEQ ID NO:9:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 615 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

Met	Trp	Arg	Gln	Ser	Thr	Ile	Leu	Ala	Ala	Leu	Leu	Val	Ala	Leu	Leu
1				5					10					15	
Cys	Ala	Gly	Ser	Ala	Glu	Ser	Lys	Gly	Asn	Arg	Pro	Pro	Arg	Ile	Thr
			20					25					30		
Lys	Gln	Pro	Ala	Pro	Gly	Glu	Leu	Phe	Lys	Val	Ala	Gln	Gln	Asn	
		35					40				45				
Lys	Glu	Ser	Asp	Pro	Glu	Arg	Asn	Pro	Phe	Ile	Ile	Glu	Cys	Glu	Ala
	50					55					60				
Asp	Gly	Gln	Pro	Glu	Pro	Glu	Tyr	Ser	Trp	Ile	Lys	Asn	Gly	Lys	Lys
	65				70					75					80
Phe	Asp	Trp	Gln	Ala	Tyr	Asp	Asn	Arg	Met	Leu	Arg	Gln	Pro	Gly	Arg
			85						90					95	
Gly	Thr	Leu	Val	Ile	Thr	Ile	Pro	Lys	Asp	Glu	Asp	Arg	Gly	His	Tyr
			100					105					110		
Gln	Cys	Phe	Ala	Ser	Asn	Glu	Phe	Gly	Thr	Ala	Thr	Ser	Asn	Ser	Val
		115					120					125			
Tyr	Val	Arg	Lys	Ala	Glu	Leu	Asn	Ala	Phe	Lys	Asp	Glu	Ala	Ala	Lys
	130					135					140				
Thr	Leu	Glu	Ala	Val	Glu	Gly	Glu	Pro	Phe	Met	Leu	Lys	Cys	Ala	Ala
	145				150					155					160
Pro	Asp	Gly	Phe	Pro	Ser	Pro	Thr	Val	Asn	Trp	Met	Ile	Gln	Glu	Ser
				165					170					175	
Ile	Asp	Gly	Ser	Ile	Lys	Ser	Ile	Asn	Asn	Ser	Arg	Met	Thr	Leu	Asp
			180					185					190		
Pro	Glu	Gly	Asn	Leu	Trp	Phe	Ser	Asn	Val	Thr	Arg	Glu	Asp	Ala	Ser
		195					200					205			
Ser	Asp	Phe	Tyr	Tyr	Ala	Cys	Ser	Ala	Thr	Ser	Val	Phe	Arg	Ser	Glu
	210					215					220				
Tyr	Lys	Ile	Gly	Asn	Lys	Val	Leu	Leu	Asp	Val	Lys	Gln	Met	Gly	Val
	225				230					235					240
Ser	Ala	Ser	Gln	Asn	Lys	His	Pro	Pro	Val	Arg	Gln	Tyr	Val	Ser	Arg
				245					250					255	
Arg	Gln	Ser	Ala	Leu	Arg	Gly	Lys	Arg	Met	Glu	Leu	Phe	Cys	Ile	Tyr
			260					265					270		
Gly	Gly	Thr	Pro	Leu	Pro	Gln	Thr	Val	Trp	Ser	Lys	Asp	Gly	Gln	Arg
		275					280					285			
Ile	Gln	Trp	Ser	Asp	Arg	Ile	Thr	Gln	Gly	His	Tyr	Gly	Lys	Ser	Leu
	290					295					300				
Val	Ile	Arg	Gln	Thr	Asn	Phe	Asp	Asp	Ala	Gly	Thr	Tyr	Thr	Cys	Asp
	305				310					315					320
Val	Ser	Asn	Gly	Val	Gly	Asn	Ala	Gln	Ser	Phe	Ser	Ile	Ile	Leu	Asn
			325					330						335	
Val	Asn	Ser	Val	Pro	Tyr	Phe	Thr	Lys	Glu	Pro	Glu	Ile	Ala	Thr	Ala
			340					345					350		
Ala	Glu	Asp	Glu	Glu	Val	Val	Phe	Glu	Cys	Arg	Ala	Ala	Gly	Val	Pro
		355					360					365			
Glu	Pro	Lys	Ile	Ser	Trp	Ile	His	Asn	Gly	Lys	Pro	Ile	Glu	Gln	Ser
	370					375					380				
Thr	Pro	Asn	Pro	Arg	Arg	Thr	Val	Thr	Asp	Asn	Thr	Ile	Arg	Ile	Ile
	385				390					395					400
Asn	Leu	Val	Lys	Gly	Asp	Thr	Gly	Asn	Tyr	Gly	Cys	Asn	Ala	Thr	Asn
			405						410					415	
Ser	Leu	Gly	Tyr	Val	Tyr	Lys	Asp	Val	Tyr	Leu	Asn	Val	Gln	Ala	Glu
		420						425					430		
Pro	Pro	Thr	Ile	Ser	Glu	Ala	Pro	Ala	Ala	Val	Ser	Thr	Val	Asp	Gly
		435					440					445			
Arg	Asn	Val	Thr	Ile	Lys	Cys	Arg	Val	Asn	Gly	Ser	Pro	Lys	Pro	Leu
	450					455					460				
Val	Lys	Trp	Leu	Arg	Ala	Ser	Asn	Trp	Leu	Thr	Gly	Gly	Arg	Tyr	Asn
	465				470					475					480
Val	Gln	Ala	Asn	Gly	Asp	Leu	Glu	Ile	Gln	Asp	Val	Thr	Phe	Ser	Asp
			485						490					495	
Ala	Gly	Lys	Tyr	Thr	Cys	Tyr	Ala	Gln	Asn	Lys	Phe	Gly	Glu	Ile	Gln
			500					505					510		
Ala	Asp	Gly	Ser	Leu	Val	Val	Lys	Glu	His	Thr	Ile	Thr	Gln	Glu	Pro
		515					520					525			

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Gln Asn Tyr Glu Val Ala Ala Gly Gln Ser Ala Thr Phe Arg Cys Asn  
 530 535 540  
 Glu Ala His Asp Asp Thr Leu Glu Ile Glu Ile Asp Trp Trp Lys Asp  
 545 550 555 560  
 Gly Gln Ser Ile Asp Phe Glu Ala Gln Pro Arg Phe Val Lys Thr Asn  
 565 570 575  
 Asp Asn Ser Leu Thr Ile Ala Lys Thr Met Glu Leu Asp Ser Gly Glu  
 580 585 590  
 Tyr Thr Cys Val Ala Arg Thr Arg Leu Asp Glu Ala Thr Ala Arg Ala  
 595 600 605  
 Asn Leu Ile Val Gln Asp Val  
 610 615

## (2) INFORMATION FOR SEQ ID NO:10:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 611 amino acids  
 (B) TYPE: amino acid  
 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

Met Val Val Ala Leu Arg Tyr Val Trp Pro Leu Leu Leu Cys Ser Pro  
 1 5 10 15  
 Cys Leu Leu Ile Gln Ile Pro Glu Glu Tyr Glu Gly His His Val Met  
 20 25 30  
 Glu Pro Pro Val Ile Thr Glu Gln Ser Pro Arg Arg Leu Val Val Phe  
 35 40 45  
 Pro Thr Asp Asp Ile Ser Leu Lys Cys Glu Ala Ser Gly Lys Pro Glu  
 50 55 60  
 Val Gln Phe Arg Trp Thr Arg Asp Gly Val His Phe Lys Pro Lys Glu  
 65 70 75 80  
 Glu Leu Gly Val Thr Val Tyr Gln Ser Pro His Ser Gly Ser Phe Thr  
 85 90 95  
 Ile Thr Gly Asn Asn Ser Asn Phe Ala Gln Arg Phe Gln Gly Ile Tyr  
 100 105 110  
 Arg Cys Phe Ala Ser Asn Lys Leu Gly Thr Ala Met Ser His Glu Ile  
 115 120 125  
 Arg Leu Met Ala Glu Gly Ala Pro Lys Trp Pro Lys Glu Thr Val Lys  
 130 135 140  
 Pro Val Glu Val Glu Glu Glu Ser Val Val Leu Pro Cys Asn Pro  
 145 150 155 160  
 Pro Pro Ser Ala Glu Pro Leu Arg Ile Tyr Trp Met Asn Ser Lys Ile  
 165 170 175  
 Leu His Ile Lys Gln Asp Glu Arg Val Thr Met Gly Gln Asn Gly Asn  
 180 185 190  
 Leu Tyr Phe Ala Asn Val Leu Thr Ser Asp Asn His Ser Asp Tyr Ile  
 195 200 205  
 Cys His Ala His Phe Pro Gly Thr Arg Thr Ile Ile Gln Lys Glu Pro  
 210 215 220  
 Ile Asp Leu Arg Val Lys Ala Thr Asn Ser Met Ile Asp Arg Lys Pro  
 225 230 235 240  
 Arg Leu Leu Phe Pro Thr Asn Ser Ser Ser His Leu Val Ala Leu Gln  
 245 250 255  
 Gly Gln Pro Leu Val Leu Glu Cys Ile Ala Glu Gly Phe Pro Thr Pro  
 260 265 270  
 Thr Ile Lys Trp Leu Arg Pro Ser Gly Pro Met Pro Ala Asp Arg Val  
 275 280 285  
 Thr Tyr Gln Asn His Asn Lys Thr Leu Gln Leu Leu Lys Val Gly Glu  
 290 295 300  
 Glu Asp Asp Gly Glu Tyr Arg Cys Leu Ala Glu Asn Ser Leu Gly Ser  
 305 310 315 320  
 Ala Arg His Ala Tyr Tyr Val Thr Val Glu Ala Ala Lys Tyr Arg Ile  
 325 330 335  
 Gln Arg Gly Ala Leu Ile Leu Ser Asn Val Gln Pro Ser Asp Thr Met  
 340 345 350



```

Val Thr Gln Cys Glu Ala Arg Asn Arg His Gly Leu Leu Leu Ala Asn
      355      360      365
Ala Tyr Ile Tyr Val Val Gln Leu Pro Ala Lys Ile Leu Thr Ala Asp
      370      375      380
Asn Gln Thr Tyr Met Ala Val Pro Tyr Trp Leu His Lys Pro Gln Ser
      385      390      395      400
His Leu Tyr Gly Pro Gly Glu Thr Ala Arg Leu Asp Cys Gln Val Gln
      405      410      415
Gly Arg Pro Gln Pro Glu Val Thr Trp Arg Ile Asn Gly Ile Pro Val
      420      425      430
Glu Glu Leu Ala Lys Asp Gln Gln Gly Ser Thr Ala Tyr Leu Leu Cys
      435      440      445
Lys Ala Phe Gly Ala Pro Val Pro Ser Val Gln Trp Leu Asp Glu Asp
      450      455      460
Gly Thr Thr Val Leu Gln Asp Glu Arg Phe Phe Pro Tyr Ala Asn Gly
      465      470      475      480
Thr Leu Gly Ile Arg Asp Leu Gln Ala Asn Asp Thr Gly Arg Tyr Phe
      485      490      495
Cys Leu Ala Ala Asn Asp Gln Asn Asn Val Thr Ile Met Ala Asn Leu
      500      505      510
Lys Val Lys Asp Ala Thr Gln Ile Thr Gln Gly Pro Arg Ser Thr Ile
      515      520      525
Glu Lys Lys Gly Ser Arg Val Thr Phe Thr Cys Gln Ala Ser Phe Asp
      530      535      540
Pro Ser Leu Gln Pro Ser Ile Thr Trp Arg Gly Asp Gly Arg Asp Leu
      545      550      555      560
Gln Glu Leu Gly Asp Ser Asp Lys Tyr Phe Ile Glu Asp Gly Arg Leu
      565      570      575
Val Ile His Ser Leu Asp Tyr Ser Asp Gln Gly Asn Tyr Ser Cys Val
      580      585      590
Ala Ser Thr Glu Leu Asp Val Val Glu Ser Arg Ala Gln Leu Leu Val
      595      600      605
Val Gly Ser
      610

```

## (2) INFORMATION FOR SEQ ID NO:11:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 612 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:11:

```

Met Met Lys Glu Lys Ser Ile Ser Ala Ser Lys Ala Ser Leu Val Phe
  1      5      10      15
Phe Leu Cys Gln Met Ile Ser Ala Leu Asp Val Pro Leu Asp Ser Lys
      20      25      30
Leu Leu Glu Glu Leu Ser Gln Pro Pro Thr Ile Thr Gln Gln Ser Pro
      35      40      45
Lys Asp Tyr Ile Val Asp Pro Arg Glu Asn Ile Val Ile Gln Cys Glu
      50      55      60
Ala Lys Gly Lys Pro Pro Pro Ser Phe Ser Trp Thr Arg Asn Gly Thr
      65      70      75      80
His Phe Asp Ile Asp Lys Asp Ala Gln Val Thr Met Lys Pro Asn Ser
      85      90      95
Gly Thr Leu Val Val Asn Ile Met Asn Gly Val Lys Ala Glu Ala Tyr
      100      105      110
Glu Gly Val Tyr Gln Cys Thr Ala Arg Asn Glu Arg Gly Ala Ala Ile
      115      120      125
Ser Asn Asn Ile Val Ile Arg Pro Ser Arg Ser Pro Leu Trp Thr Lys
      130      135      140
Glu Lys Leu Glu Pro Asn His Val Arg Glu Gly Asp Ser Leu Val Leu
      145      150      155      160
Asn Cys Arg Pro Pro Val Gly Leu Pro Pro Pro Ile Ile Phe Trp Met
      165      170      175

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Asp Asn Ala Phe Gln Arg Leu Pro Gln Ser Glu Arg Val Ser Gln Gly
      180      185      190
Leu Asn Gly Asp Leu Tyr Phe Ser Asn Val Gln Pro Glu Asp Thr Arg
      195      200      205
Val Asp Tyr Ile Cys Tyr Ala Arg Phe Asn His Thr Gln Thr Ile Gln
      210      215      220
Gln Lys Gln Pro Ile Ser Val Lys Val Phe Ser Thr Lys Pro Val Thr
225      230      235      240
Glu Arg Pro Pro Val Leu Leu Thr Pro Met Gly Ser Thr Ser Asn Lys
      245      250      255
Val Glu Leu Arg Gly Asn Val Leu Leu Leu Glu Cys Ile Ala Ala Gly
      260      265      270
Leu Pro Thr Pro Val Ile Arg Trp Ile Lys Glu Gly Gly Glu Leu Pro
      275      280      285
Ala Asn Arg Thr Phe Phe Glu Asn Phe Lys Lys Thr Leu Lys Ile Ile
      290      295      300
Asp Val Ser Glu Ala Asp Ser Gly Asn Tyr Lys Cys Thr Ala Arg Asn
305      310      315      320
Thr Leu Gly Ser Thr His His Val Ile Ser Val Thr Val Lys Ala Ala
      325      330      335
Pro Tyr Trp Ile Thr Ala Pro Arg Asn Leu Val Leu Ser Pro Gly Glu
      340      345      350
Asp Gly Thr Leu Ile Cys Arg Ala Asn Gly Asn Pro Lys Pro Ser Ile
      355      360      365
Ser Trp Leu Thr Asn Gly Val Pro Ile Ala Ile Ala Pro Glu Asp Pro
      370      375      380
Ser Arg Lys Val Asp Gly Asp Thr Ile Ile Phe Ser Ala Val Gln Glu
385      390      395      400
Arg Ser Ser Ala Val Tyr Gln Cys Asn Ala Ser Asn Glu Tyr Gly Tyr
      405      410      415
Leu Leu Ala Asn Ala Phe Val Asn Val Leu Ala Glu Pro Pro Arg Ile
      420      425      430
Leu Thr Pro Ala Asn Lys Leu Tyr Gln Val Ile Ala Asp Ser Pro Ala
      435      440      445
Leu Ile Asp Cys Ala Tyr Phe Gly Ser Pro Lys Pro Glu Ile Glu Trp
      450      455      460
Phe Arg Gly Val Lys Gly Ser Ile Leu Arg Gly Asn Glu Tyr Val Phe
465      470      475      480
His Asp Asn Gly Thr Leu Glu Ile Pro Val Ala Gln Lys Asp Ser Thr
      485      490      495
Gly Thr Tyr Thr Cys Val Ala Arg Asn Lys Leu Gly Lys Thr Gln Asn
      500      505      510
Glu Val Gln Leu Glu Val Lys Asp Pro Thr Met Ile Ile Lys Gln Pro
      515      520      525
Gln Tyr Lys Val Ile Gln Arg Ser Ala Gln Ala Ser Phe Glu Cys Val
      530      535      540
Ile Lys His Asp Pro Thr Leu Ile Pro Thr Val Ile Trp Leu Lys Asp
545      550      555      560
Asn Asn Glu Leu Pro Asp Asp Glu Arg Phe Leu Val Gly Lys Asp Asn
      565      570      575
Leu Thr Ile Met Asn Val Thr Asp Lys Asp Asp Gly Thr Tyr Thr Cys
      580      585      590
Ile Val Asn Thr Thr Leu Asp Ser Val Ser Ala Ser Ala Val Leu Thr
      595      600      605
Val Val Ala Ala
610

```

## (2) INFORMATION FOR SEQ ID NO:12:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 607 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:12:

Met	Gly	Thr	Ala	Thr	Arg	Arg	Lys	Pro	His	Leu	Leu	Leu	Val	Ala	Ala	1	5	10	15
Val	Ala	Leu	Val	Ser	Ser	Ser	Ala	Trp	Ser	Ser	Ala	Leu	Gly	Ser	Gln	20	25	30	
Thr	Thr	Phe	Gly	Pro	Val	Phe	Glu	Asp	Gln	Pro	Leu	Ser	Val	Leu	Phe	35	40	45	
Pro	Glu	Glu	Ser	Thr	Glu	Glu	Gln	Val	Leu	Leu	Ala	Cys	Arg	Ala	Arg	50	55	60	
Ala	Ser	Pro	Pro	Ala	Thr	Tyr	Arg	Trp	Lys	Met	Asn	Gly	Thr	Glu	Met	65	70	75	80
Lys	Leu	Glu	Pro	Gly	Ser	Arg	His	Gln	Leu	Val	Gly	Gly	Asn	Leu	Val	85	90	95	
Ile	Met	Asn	Pro	Thr	Lys	Ala	Gln	Asp	Ala	Gly	Val	Tyr	Gln	Cys	Leu	100	105	110	
Ala	Ser	Asn	Pro	Val	Gly	Thr	Val	Ser	Arg	Glu	Ala	Ile	Leu	Arg		115	120	125	
Phe	Gly	Phe	Leu	Gln	Glu	Phe	Ser	Lys	Glu	Glu	Arg	Asp	Pro	Val	Lys	130	135	140	
Ala	His	Glu	Gly	Trp	Gly	Val	Met	Leu	Pro	Cys	Asn	Pro	Pro	Ala	His	145	150	155	160
Tyr	Pro	Gly	Leu	Ser	Tyr	Arg	Trp	Leu	Leu	Asn	Glu	Phe	Pro	Asn	Phe	165	170	175	
Ile	Pro	Thr	Asp	Gly	Arg	His	Phe	Val	Ser	Gln	Thr	Thr	Gly	Asn	Leu	180	185	190	
Tyr	Ile	Ala	Arg	Thr	Asn	Ala	Ser	Asp	Leu	Gly	Asn	Tyr	Ser	Cys	Leu	195	200	205	
Ala	Thr	Ser	His	Met	Asp	Phe	Ser	Thr	Lys	Ser	Val	Phe	Ser	Lys	Phe	210	215	220	
Ala	Gln	Leu	Asn	Leu	Ala	Ala	Glu	Asp	Thr	Arg	Leu	Phe	Ala	Pro	Ser	225	230	235	240
Ile	Lys	Ala	Arg	Phe	Pro	Ala	Glu	Thr	Tyr	Ala	Leu	Val	Gly	Gln	Gln	245	250	255	
Val	Thr	Leu	Glu	Cys	Phe	Ala	Phe	Gly	Asn	Pro	Val	Pro	Arg	Ile	Lys	260	265	270	
Trp	Arg	Lys	Val	Asp	Gly	Ser	Leu	Ser	Pro	Gln	Trp	Thr	Thr	Ala	Glu	275	280	285	
Pro	Thr	Leu	Gln	Ile	Pro	Ser	Val	Ser	Phe	Glu	Asp	Glu	Gly	Thr	Tyr	290	295	300	
Glu	Cys	Glu	Ala	Glu	Asn	Ser	Lys	Gly	Arg	Asp	Thr	Val	Gln	Gly	Arg	305	310	315	320
Ile	Ile	Val	Gln	Ala	Gln	Pro	Glu	Trp	Leu	Lys	Val	Ile	Ser	Asp	Thr	325	330	335	
Glu	Ala	Asp	Ile	Gly	Ser	Asn	Leu	Arg	Trp	Gly	Cys	Ala	Ala	Ala	Gly	340	345	350	
Lys	Pro	Arg	Pro	Thr	Val	Arg	Trp	Leu	Arg	Asn	Gly	Glu	Pro	Leu	Ala	355	360	365	
Ser	Gln	Asn	Arg	Val	Glu	Val	Leu	Ala	Gly	Asp	Leu	Arg	Phe	Ser	Lys	370	375	380	
Leu	Ser	Leu	Glu	Asp	Ser	Gly	Met	Tyr	Gln	Cys	Val	Ala	Glu	Asn	Lys	385	390	395	400
His	Gly	Thr	Ile	Tyr	Ala	Ser	Ala	Glu	Leu	Ala	Val	Gln	Ala	Leu	Ala	405	410	415	
Pro	Asp	Phe	Arg	Leu	Asn	Pro	Val	Arg	Arg	Leu	Ile	Pro	Ala	Ala	Arg	420	425	430	
Gly	Gly	Glu	Ile	Leu	Ile	Pro	Cys	Gln	Pro	Arg	Ala	Ala	Pro	Lys	Ala	435	440	445	
Val	Val	Leu	Trp	Ser	Lys	Gly	Thr	Glu	Ile	Leu	Val	Asn	Ser	Ser	Arg	450	455	460	
Val	Thr	Val	Thr	Pro	Asp	Gly	Thr	Leu	Ile	Ile	Arg	Asn	Ile	Ser	Arg	465	470	475	480
Ser	Asp	Glu	Gly	Lys	Tyr	Thr	Cys	Phe	Ala	Glu	Asn	Phe	Met	Gly	Lys	485	490	495	
Ala	Asn	Ser	Thr	Gly	Ile	Leu	Ser	Val	Arg	Asp	Ala	Thr	Lys	Ile	Thr	500	505	510	
Leu	Ala	Pro	Ser	Ser	Ala	Asp	Ile	Asn	Leu	Gly	Asp	Asn	Leu	Thr	Leu	515	520	525	

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Gln	Cys	His	Ala	Ser	His	Asp	Pro	Thr	Met	Asp	Leu	Thr	Phe	Thr	Trp
	530					535					540				
Thr	Leu	Asp	Asp	Phe	Pro	Ile	Asp	Phe	Asp	Lys	Pro	Gly	Gly	His	Tyr
545					550					555					560
Arg	Arg	Thr	Asn	Val	Lys	Glu	Thr	Ile	Gly	Asp	Leu	Thr	Ile	Leu	Asn
			565						570					575	
Ala	Gln	Leu	Arg	His	Gly	Gly	Lys	Tyr	Thr	Cys	Met	Ala	Gln	Thr	Val
			580					585					590		
Val	Asp	Ser	Ala	Ser	Lys	Glu	Ala	Thr	Val	Leu	Val	Arg	Gly	Pro	
	595						600					605			

## (2) INFORMATION FOR SEQ ID NO:13:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 596 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:13:

Met	Leu	Ser	Trp	Lys	Gln	Leu	Ile	Leu	Leu	Ser	Phe	Ile	Gly	Cys	Leu
1				5				10					15		
Ala	Gly	Glu	Leu	Leu	Gln	Gly	Pro	Val	Phe	Val	Lys	Glu	Pro	Ser	
			20				25					30			
Asn	Ser	Ile	Phe	Pro	Val	Gly	Ser	Glu	Asp	Lys	Lys	Ile	Thr	Leu	Asn
		35				40						45			
Cys	Glu	Ala	Arg	Gly	Asn	Pro	Ser	Pro	His	Tyr	Arg	Trp	Gln	Leu	Asn
	50				55						60				
Gly	Ser	Asp	Ile	Asp	Thr	Ser	Leu	Asp	His	Arg	Tyr	Lys	Leu	Asn	Gly
65				70					75					80	
Gly	Asn	Leu	Ile	Val	Ile	Asn	Pro	Asn	Arg	Asn	Trp	Asp	Thr	Gly	Ser
			85					90						95	
Tyr	Gln	Cys	Phe	Ala	Thr	Asn	Ser	Leu	Gly	Thr	Ile	Val	Ser	Arg	Glu
			100					105					110		
Ala	Lys	Leu	Gln	Phe	Ala	Tyr	Leu	Glu	Asn	Phe	Lys	Ser	Arg	Met	Arg
	115						120					125			
Ser	Arg	Val	Ser	Val	Arg	Glu	Gly	Gln	Gly	Val	Val	Leu	Leu	Cys	Gly
	130					135					140				
Pro	Pro	Pro	His	Ser	Gly	Glu	Leu	Ser	Tyr	Ala	Trp	Val	Phe	Asn	Glu
145					150					155				160	
Tyr	Pro	Ser	Phe	Val	Glu	Glu	Asp	Ser	Arg	Arg	Phe	Val	Ser	Gln	Glu
			165						170					175	
Thr	Gly	His	Leu	Tyr	Ile	Ala	Lys	Val	Glu	Pro	Ser	Asp	Val	Gly	Asn
	180							185					190		
Tyr	Thr	Cys	Val	Val	Thr	Ser	Thr	Val	Thr	Asn	Ala	Arg	Val	Leu	Gly
	195						200					205			
Ser	Pro	Thr	Pro	Leu	Val	Leu	Arg	Ser	Asp	Gly	Val	Met	Gly	Glu	Tyr
	210					215					220				
Glu	Pro	Lys	Ile	Glu	Leu	Gln	Phe	Pro	Glu	Thr	Leu	Pro	Ala	Ala	Lys
225					230					235				240	
Gly	Ser	Thr	Val	Lys	Leu	Glu	Cys	Phe	Ala	Leu	Gly	Asn	Pro	Val	Pro
			245						250					255	
Gln	Ile	Asn	Trp	Arg	Arg	Ser	Asp	Gly	Met	Pro	Phe	Pro	Thr	Lys	Ile
		260						265					270		
Lys	Leu	Arg	Lys	Phe	Asn	Gly	Val	Leu	Glu	Ile	Pro	Asn	Phe	Gln	Gln
	275						280					285			
Glu	Asp	Thr	Gly	Ser	Tyr	Glu	Cys	Ile	Ala	Glu	Asn	Ser	Arg	Gly	Lys
	290					295					300				
Asn	Val	Ala	Arg	Gly	Arg	Leu	Thr	Tyr	Tyr	Ala	Lys	Pro	Tyr	Trp	Val
305					310					315				320	
Gln	Leu	Leu	Lys	Asp	Val	Glu	Thr	Ala	Val	Glu	Asp	Ser	Leu	Tyr	Trp
			325						330					335	
Glu	Cys	Arg	Ala	Ser	Gly	Lys	Pro	Lys	Pro	Ser	Tyr	Arg	Trp	Leu	Lys
		340						345					350		
Asn	Gly	Asp	Ala	Leu	Val	Leu	Glu	Glu	Arg	Ile	Gln	Ile	Glu	Asn	Gly
	355						360					365			

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Ala Leu Thr Ile Ala Asn Leu Asn Val Ser Asp Ser Gly Met Phe Gln  
 370 375 380  
 Cys Ile Ala Glu Asn Lys His Gly Leu Ile Tyr Ser Ser Ala Glu Leu  
 385 390 395 400  
 Lys Val Leu Ala Ser Ala Pro Asp Phe Ser Arg Asn Pro Met Lys Lys  
 405 410 415  
 Met Ile Gln Val Gln Val Gly Ser Leu Val Ile Leu Asp Cys Lys Pro  
 420 425 430  
 Ser Ala Ser Pro Arg Ala Leu Ser Phe Trp Lys Lys Gly Asp Thr Val  
 435 440 445  
 Val Arg Glu Gln Ala Arg Ile Ser Leu Leu Asn Asp Gly Gly Leu Lys  
 450 455 460  
 Ile Met Asn Val Thr Lys Ala Asp Ala Gly Ile Tyr Thr Cys Ile Ala  
 465 470 475 480  
 Glu Asn Gln Phe Gly Lys Ala Asn Gly Thr Thr Gln Leu Val Val Thr  
 485 490 495  
 Glu Pro Thr Arg Ile Ile Leu Ala Pro Ser Asn Met Asp Val Ala Val  
 500 505 510  
 Gly Glu Ser Ile Ile Leu Pro Cys Gln Val Gln His Asp Pro Leu Leu  
 515 520 525  
 Asp Ile Met Phe Ala Trp Tyr Phe Asn Gly Thr Leu Thr Asp Phe Lys  
 530 535 540  
 Lys Asp Gly Ser His Phe Glu Lys Val Gly Gly Ser Ser Ser Gly Asp  
 545 550 555 560  
 Leu Met Ile Arg Asn Ile Gln Leu Lys His Ser Gly Lys Tyr Val Cys  
 565 570 575  
 Met Val Gln Thr Gly Val Asp Ser Val Ser Ser Ala Ala Glu Leu Ile  
 580 585 590  
 Val Arg Gly Ser  
 595

## (2) INFORMATION FOR SEQ ID NO:14:

## (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 630 amino acids  
 (B) TYPE: amino acid  
 (D) TOPOLOGY: linear

## (ii) MOLECULE TYPE: protein

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:14:

Met Val Leu His Ser His Gln Leu Thr Tyr Ala Gly Ile Ala Phe Ala  
 1 5 10 15  
 Leu Cys Leu His His Leu Ile Ser Ala Ile Glu Val Pro Leu Asp Ser  
 20 25 30  
 Asn Ile Gln Ser Glu Leu Pro Gln Pro Pro Thr Ile Thr Lys Gln Ser  
 35 40 45  
 Val Lys Asp Tyr Ile Val Asp Pro Arg Asp Asn Ile Phe Ile Glu Cys  
 50 55 60  
 Glu Ala Lys Gly Asn Pro Val Pro Thr Phe Ser Trp Thr Arg Asn Gly  
 65 70 75 80  
 Lys Phe Phe Asn Val Ala Lys Asp Pro Lys Val Ser Met Arg Arg Arg  
 85 90 95  
 Ser Gly Thr Leu Val Ile Asp Phe His Gly Gly Gly Arg Pro Asp Asp  
 100 105 110  
 Tyr Glu Gly Glu Tyr Gln Cys Phe Ala Arg Asn Asp Tyr Gly Thr Ala  
 115 120 125  
 Leu Ser Ser Lys Ile His Leu Gln Val Ser Arg Ser Pro Leu Trp Pro  
 130 135 140  
 Lys Glu Lys Val Asp Val Ile Glu Val Asp Glu Gly Ala Pro Leu Ser  
 145 150 155 160  
 Leu Gln Cys Asn Pro Pro Pro Gly Leu Pro Pro Pro Val Ile Phe Trp  
 165 170 175  
 Met Ser Ser Ser Met Glu Pro Ile His Gln Asp Lys Arg Val Ser Gln  
 180 185 190  
 Gly Gln Asn Gly Asp Leu Tyr Phe Ser Asn Val Met Leu Gln Asp Ala  
 195 200 205

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Gln	Thr	Asp	Tyr	Ser	Cys	Asn	Ala	Arg	Phe	His	Phe	Thr	His	Thr	Ile
	210					215					220				
Gln	Gln	Lys	Asn	Pro	Tyr	Thr	Leu	Lys	Val	Lys	Thr	Lys	Lys	Pro	His
225					230					235					240
Asn	Glu	Thr	Ser	Leu	Arg	Asn	His	Thr	Asp	Met	Tyr	Ser	Ala	Arg	Gly
				245					250					255	
Val	Thr	Glu	Thr	Thr	Pro	Ser	Phe	Met	Tyr	Pro	Tyr	Gly	Thr	Ser	Ser
			260					265					270		
Ser	Gln	Met	Val	Leu	Arg	Gly	Val	Asp	Leu	Leu	Leu	Glu	Cys	Ile	Ala
		275					280					285			
Ser	Gly	Val	Pro	Ala	Pro	Asp	Ile	Met	Trp	Tyr	Lys	Lys	Gly	Gly	Glu
	290					295					300				
Leu	Pro	Ala	Gly	Lys	Thr	Lys	Leu	Glu	Asn	Phe	Asn	Lys	Ala	Leu	Arg
305					310					315					320
Ile	Ser	Asn	Val	Ser	Glu	Glu	Asp	Ser	Gly	Glu	Tyr	Phe	Cys	Leu	Ala
				325					330					335	
Ser	Asn	Lys	Met	Gly	Ser	Ile	Arg	His	Thr	Ile	Ser	Val	Arg	Val	Lys
			340					345					350		
Ala	Ala	Pro	Tyr	Trp	Leu	Asp	Glu	Pro	Gln	Asn	Leu	Ile	Leu	Ala	Pro
		355					360					365			
Gly	Glu	Asp	Gly	Arg	Leu	Val	Cys	Arg	Ala	Asn	Gly	Asn	Pro	Lys	Pro
	370					375					380				
Ser	Ile	Gln	Trp	Leu	Val	Asn	Gly	Glu	Pro	Ile	Glu	Gly	Ser	Pro	Pro
385					390					395					400
Asn	Pro	Ser	Arg	Glu	Val	Ala	Gly	Asp	Thr	Ile	Val	Phe	Arg	Asp	Thr
				405					410					415	
Gln	Ile	Gly	Ser	Ala	Val	Tyr	Gln	Cys	Asn	Ala	Ser	Asn	Glu	His	
			420				425						430		
Gly	Tyr	Leu	Leu	Ala	Asn	Ala	Phe	Val	Ser	Val	Leu	Asp	Val	Pro	Pro
		435					440					445			
Arg	Ile	Leu	Ala	Pro	Arg	Asn	Gln	Leu	Ile	Lys	Val	Ile	Gln	Tyr	Asn
	450					455					460				
Arg	Thr	Arg	Leu	Asp	Cys	Pro	Phe	Phe	Gly	Ser	Pro	Ile	Pro	Thr	Leu
465					470					475					480
Arg	Trp	Phe	Lys	Asn	Gly	Gln	Gly	Asn	Met	Leu	Asp	Gly	Gly	Asn	Tyr
				485					490					495	
Lys	Ala	His	Glu	Asn	Gly	Ser	Leu	Glu	Met	Ser	Met	Ala	Arg	Lys	Glu
			500					505					510		
Asp	Gln	Gly	Ile	Tyr	Thr	Cys	Val	Ala	Thr	Asn	Ile	Leu	Gly	Lys	Val
		515					520					525			
Glu	Ala	Gln	Val	Arg	Leu	Glu	Val	Lys	Asp	Pro	Thr	Arg	Ile	Val	Arg
					535						540				
Gly	Pro	Glu	Asp	Gln	Val	Val	Lys	Arg	Gly	Ser	Met	Pro	Arg	Leu	His
545					550					555					560
Cys	Arg	Val	Lys	His	Asp	Pro	Thr	Leu	Lys	Leu	Thr	Val	Thr	Trp	Leu
				565					570					575	
Lys	Asp	Asp	Ala	Pro	Leu	Tyr	Ile	Gly	Asn	Arg	Met	Lys	Lys	Glu	Asp
			580					585					590		
Asp	Gly	Leu	Thr	Ile	Tyr	Gly	Val	Ala	Glu	Lys	Asp	Gln	Gly	Asp	Tyr
		595					600					605			
Thr	Cys	Val	Ala	Ser	Thr	Glu	Leu	Asp	Lys	Asp	Ser	Ala	Lys	Ala	Tyr
	610					615					620				
Leu	Thr	Val	Leu	Ala	Ile										
625					630										

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What is claimed is:

1. A method for identifying a cDNA nucleic acid encoding a mammalian protein having a signal sequence, the method comprising:

- 5           a) providing library of mammalian cDNA;
- b) ligating said library of mammalian cDNA to DNA encoding alkaline phosphatase lacking both a signal sequence and a membrane anchor sequence to form ligated DNA;
- 10          c) transforming bacterial cells with said ligated DNA to create a bacterial cell clone library;
- d) isolating DNA comprising said mammalian cDNA from at least one clone in said bacterial cell clone library;
- 15          e) separately transfecting DNA isolated from clones in step (d) into mammalian cells which do not express alkaline phosphatase to create a mammalian cell clone library wherein each clone in said mammalian cell clone library corresponds to a clone in said bacterial
- 20 cell clone library;
- f) identifying a clone in said mammalian cell clone library which express alkaline phosphatase;
- g) identifying the clone in said bacterial cell clone library corresponding to said clone in said
- 25 mammalian cell clone library identified in step (f); and
- h) isolating and sequencing a portion of the mammalian cDNA present in said bacterial cell library clone identified in step (g) to identify a mammalian cDNA encoding a mammalian protein having a signal sequence.
- 30          2. The method of claim 1 wherein said library of mammalian cDNAs are ligated to ptrAP3.

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3. The method of claim 1 wherein said mammalian cells are COS7 cells.

4. The method of claim 1 wherein said bacterial cells are E. coli.

5 5. The expression vector ptrAP3.

6. The expression vector of claim 5, comprising the sequence of SEQ ID NO:1.

7. The protein of SEQ ID NO:5.

8. An isolated nucleic acid sequence encoding the  
10 amino acid sequence of SEQ ID NO:5.

9. A vector comprising the nucleic acid sequence of claim 8.

10. The vector of claim 9 wherein said vector is an expression vector.

15 11. A genetically engineered host cell comprising the nucleic acid sequence of claim 5.



**ptrAP3****FIG. 1**

**ptrAP3 vector sequence**

AAGCTTGGCTGTGGAATGTGTGTCAGTTAGGGTGTGGAAGTCCCCAGGCTCCCCAGCAGGCAGAAGTATGC  
AAAGCATGCATCTCAATTAGTCAGCAACCAGGTGTGGAAGTCCCCAGGCTCCCCAGCAGGCAGAAGTATGC  
AAAGCATGCATCTCAATTAGTCAGCAACCATAGTCCCGCCCCCTAACTCCGCCCCATCCGCCCCCTAACTCCGC  
CCAGTTCGCCCCATTCTCCGCCCCATGGCTGACTAATTTTTTTTATTTATGCAGAGGCCGAGGCCGCCCTCGG  
CCTCTGAGCTATTCCAGAAGTAGTGAGGAGGCTTTTTTGGAGGCCCTAGGCTTTTGCAAAAAGCTCCTCCGAT  
CGAGGGGCTCGCATCTCTCCTTCACGCGCCCGCCGCCCTACCTGAGGCCGCCATCCACGCCGGTTGAGTCGC  
GTTCTGCCGCCCTCCCGCCTGTGGTGCCTCCTGAACTGCGTCCGCCGTCTAGGTAAGTTTAAAGCTCAGGTCCG  
AGACCGGGCCTTTGTCCGGCGCTCCCTTGAGCCTACCTAGACTCAGCCGGCTCTCCACGCTTTGCCGTGACC  
CTGCTTGCTCAACTCTACGTCTTTGTTTCGTTTTCTGTTCTGCGCCGTTACAGATCCAAGCTCTGAAAACC  
AGAAAGTTAACTGGTAAGTTTAGTCTTTTGTCTTTTATTTTCAGGTCCCAGGTCCCGGATCCGGTGATCCAA  
ATCTAAGAACTGCTCCTCAGTGAGTGTTCCTTTACTTCTAGGCCTGTACGGAAGTGTACTTCTGCTCTAA  
AAGCTGCGGAATTCGCACCCAGCTAGTTTTTACGCCCCGGTGAAGCCTCCACCCGCACCTAGA  
<sup>510</sup>  
AGCGCGTGTATGATGAGGTGTACGGCGACGAGGACCTGCTTGAAGCAGGCCAACGAGCGCCT  
CGGGGAGTTTGCTACGGAAAAGCGGCATAAGGACATGTTGGCGTTGCCGCTGGACGAGGGC  
AACCCAAACACCTAGCCTAAAGCCCCGTGACACTGCAGCAGGTGCTGCCACCGCTTGACCCGT  
CCGAAGAAAAGCGCGGCCCTAAAGCGCGAGTCTGGTGACTTGGCACCCACCGTGCAGCTGAT  
GGTACCCAAAGCGCCAGCGACTGGAAGATGCTCTTGGAAAAAATGACCGTGGAGCCTGGGCTG  
GAGCCCCGAGGTCCGCGTGCGGCCAATCAAGCAGGTGGCACCGGGAAGTGGGCGTGACAGACCG  
TGGACGTTCAAGATACCCACCACCAAGTAGCACTAGTATTGCCACTGCCACAGAGGGCATGGA  
GACACAAACGTCCCCGGTTGCCCTAGCTCGAGATCATCCCAGTTGAGGAGGAGAACCCGGACTTCTG  
GAACCGCGAGGCAGCCGAGGCCCTGGGTGCCGCCAAGAAGCTGCAGCCTGCACAGACAGCCGCCAAGAACCT  
CATCATCTTCCCTGGGCGATGGGATGGGGGTGCTCTACGGTGACAGCTGCCAGGATCCTAAAAGGGCAGAAGAA  
GGACAAACTGGGGCCTGAGATACCCCTGGCCATGGACCGCTTCCCATATGTGGCTCTGTCCAAGACATACAA  
TGTAGACAAACATGTGCCAGACAGTGGAGCCACAGCCACGGCCTACCTGTGCGGGGTCAAGGGCAACTTCCA  
GACCATTTGGCTTGAGTGCAGCCGCCCGCTTTAAACCAGTGCAACACGACAGCGGCCAACGAGGTGATCTCCGT  
GATGAATCGGGCCAAGAAAGCAGGGAAGTCAGTGGGAGTGGTAACCACACAGAGTGCAGCACGCCCTCGCC  
AGCCGGCACCTACGCCCCACCGGTGAACCGCAACTGGTACTCGGACGCCGACGTCCTGCCTCGGCCCCGCCA  
GGAGGGGTGCCAGGACATCGCTACGCAGCTCATCTCCAACATGGACATTGACGTTGATCCTAGGTGGAGGCCG

FIG. 2

AAAGTACATGTTTCGCATGGGAACCCAGACCCCTGAGTACCCAGATGACTACAGCCAAGGTGGGACCAGGCT  
GGACGGGAAGAATCTGGTGCAGGAATGGCTGGCGAAGCGCCAGGGTGCCCGGTATGTGTGGAACCGCACTGA  
GCTCATGCAGGCTTCCTCGACCCGCTCTGTGACCCATCTCATGGGTCTCTTTGAGCCTGGAGACATGAAATA  
CGAGATCCACCGAGACTCCACACTGGACCCCTCCCTGATGGAGATGACAGAGGCTGCCCTGCGCCTGCTGAG  
CAGGAACCCCCGCGGCTTCTTCCTCTTCGTGGAGGGTGGTTCGCATCGACCATGGTCATCATGAAAGCAGGGC  
TTACCGGGCACTGACTGAGACGATCATGTTTCGACGACGCCATTGAGAGGGCGGGCCAGCTCACCAGCGAGGA  
GGACACGCTGAGCCTCGTCACTGCCGACCACTCCACGCTCTTCTCCTTCGGAGGCTACCCCTGCGAGGGAG  
CTCCATCTTCGGGCTGGCCCCCTGGCAAGGCCCCGGACAGGAAGGCTACACGGTCTCTCTATACGGAAACGG  
TCCAGGCTATGTGCTCAAGGACGGCGCCCGCGGATGTTACCGAGAGCGAGAGCGGGAGCCCCGAGTATCG  
GCAGCAGTCAGCAGTGGCCCTGGACGAAGAGACCCACGCAGGCGAGGACGTGGCGGTGTTTCGCGCGCGCCCC  
GCAGGCGCACCTGGTTACGGCGTGCAGGAGCAGACCTTCATAGCGCACGTATGGCCTTCGCCGCTGCGCT  
GGAGCCCTACACCGCTGCGACCTGGCGCCCCCGCGCGCACCCAGCGCGCGCACCCGGGTTGAACTAG  
TCTAGAGAAAAACCTCCACACCTCCCCCTGAACCTGAAACATAAAATGAATGCAATTGTTGTTGTTAACT  
TGTTTATTGCAGCTTATAATGGTTACAAATAAAGCAATAGCATCACAAATTCACAAATAAAGCATTTTTTTT  
CACTGCATTCTAGTTGTGTTTGTCCAACTCATCAATGTATCTTATCATGTCTGGATCCCCGGGTACCGAG  
CTCGAATTAATTCTCTTCGCTTCCTCGCTCACTGACTCGCTGCGCTCGGTCTGCTGCGCTGCGGCGAGCGG  
TATCAGCTCACTCAAAGGCGGTAATACGGTTATCCACAGAATCAGGGGATAACGCAGGAAAGAACATGTGAG  
CAAAAGGCCAGCAAAAGGCCAGGAACCGTAAAAAGGCCGCGTTGCTGGCGTTTTTCCATAGGCTCCGCCCCC  
CTGACGAGCATCACAAAAATCGACGCTCAAGTCAGAGGTGGCGAAACCCGACAGGACTATAAAGATACCAGG  
CGTTTCCCCCTGGAAGCTCCCTCGTGCGCTCTCTGTTCCGACCCTGCCGCTTACCGGATACCTGTCCGCCT  
TTCTCCCTTCGGGAAGCGTGGCGCTTTCTCAATGCTCACGCTGTAGGTATCTCAGTTCCGGTGTAGGTCTGTT  
GCTCCAAGCTGGGCTGTGTGCACGAACCCCCCGTTTCAGCCCGACCGCTGCGCCTTATCCGGTAACTATCGTC  
TTGAGTCCAACCCGGTAAGACACGACTTATCGCCACTGGCAGCAGCCACTGGTAACAGGATTAGCAGAGCGA  
GGTATGTAGGCGGTGCTACAGAGTTCTTGAAGTGGTGGCCTAACTACGGCTACACTAGAAGGACAGTATTTG  
GTATCTGCGCTCTGCTGAAGCCAGTTACCTTCGGAAAAAGAGTTGGTAGCTCTTGATCCGGCAAACAAACCA  
CCGCTGGTAGCGGTGGTTTTTTTTGTTTGCAAGCAGCAGATTACGCGCAGAAAAAAGGATCTCAAGAAGATC  
CTTTGATCTTTTCTACGGGGTCTGACGCTCAGTGAACGAAAACCTACGTTAAGGGATTTTGGTTCATGAGAT  
TATCAAAAAGGATCTTCACCTAGATCCCTTTTAAATTAAAAATGAAGTTTTAAATCAATCTAAAGTATATAG  
AGTAAACTTGGTCTGACAGTTACCAATGCTTAATCAGTGAGGCACCTATCTCAGCGATCTGTCTATTTTCGTT  
CATCCATAGTTGCCTGACTCCCCGTCGTGTAGATAACTACGATACGGGAGGGCTTACCATCTGGCCCCAGTG  
CTGCAATGATACCGCGAGACCCACGCTCACCGGCTCCAGATTTATCAGCAATAAACCAGCCAGCCGGAAGGG  
CCGAGCGCAGAAGTGGTCTGCAACTTTATCCGCCCTCCATCCAGTCTATTAATTGTTGCCGGAAGCTAGAG  
TAAGTAGTTCCGCCAGTTAATAGTTTGGCGAACGTTGTTGCCATTGCTACAGGCATCGTGGTGTACGCTCGT  
CGTTTGGTATGGCTTCATTCAGCTCCGGTTCCCAACGATCAAGGCGAGTTACATGATCCCCCATGTTGTGCA  
AAAAAGCGGTTAGCTCCTTCGGTCTCCGATCGTTGTCAGAAGTAAGTTGGCCGCAGTGTTATCACTCATGG

FIG. 2

TTATGGCAGCACTGCATAATTCTCTTACTGTCATGCCATCCGTAAGATGCTTTTCTGTGACTGGTGAGTACT  
CAACCAAGTCATTCTGAGAATAGTGTATGCGGCGACCGAGTTGCTCTTGCCCGGCGTCAATACGGGATAATA  
CCGCGCCACATAGCAGAACTTTAAAAGTGCTCATCATTGGAAAACGTTCTTCGGGGCGAAAACCTCTCAAGGA  
TCTTACCGCTGTTGAGATCCAGTTCGATGTAACCCACTCGTGCACCCCACTGATCTTCAGCATCTTTTACTT  
TCACCAGCGTTTCTGGGTGAGCAAAAACAGGAAGGCAAAATGCCGCAAAAAGGGAATAAGGGCGACACGGA  
AATGTTGAATACTCATACTCTTCCTTTTTCAATATTATTGAAGCATTTATCAGGGTTATTGTCTCATGAGCG  
GATACATATTTGAATGTATTTAGAAAAATAAACAAATAGGGGTTCGCGCACATTTCCCCGAAAAGTGCCAC  
CTGC

(SEQ ID NO: 1)  
2

FIG. 2

FIG. 3

MLLLLLLLGLRLQLSLGII PVEEENPDFWNREAAEALGAAKKLQPAQTAAKNLI  
IFLGDGMGVSTVTAARILKGQKKDKLGP EIP LAMDRFPYVALSKTYNVDKHVPD  
SGATATAYLCGVKGNFQTIGLSAAARFNQCNTTRGNEVISVMNRAKKAGKSVG  
VTTTRVQHASPAGTYAHTVNRNWYSDADVPASARQEGCQDIATQLISNMDIDVI  
LGGGRKYMFRMGTPDPEYPDDYSQGGTRLDGKNLVQEWLAKRQGARYVWNRT  
ELMQASLDPSVTHLMGLFEPGDMKYEIHRDSTLDPSLMEMTEAALRLLSRNPRGFF  
LFVEGGRIDHGHESRAYRALTETIMFDDAIERAGQLTSEEDTLSLVTADHSHV  
FSFGGYPLRGSSIFGLAPGKARDRKAYTVLLYGNGPGYVLKDGARPDVTESESG  
SPEYRQQSAVPLDEETHAGEDVAVFARGPQAHLVHGVQEQT FIAHVMAFAACLE  
PYTACDLAPPAGTTDAAHPGRSVVPALLPLLAGTLLLETTATAP

(SEE ID NO: 2)

FIG. 4

IIPVEEENPDFWNREAAEALGAAKKLQPAQTAAKNLIIFLGDGMGVSTVTAARI  
LKGQKKDKLGP EIP LAMDRFPYVALSKTYNVDKHVPD SGATATAYLCGVKGNFQ  
TIGLSAAARFNQCNTTRGNEVISVMNRAKKAGKSVG VTTTRVQHASPAGTYAH  
TVNRNWYSDADVPASARQEGCQDIATQLISNMDIDVILGGGRKYMFRMGTPDPE  
YPDDYSQGGTRLDGKNLVQEWLAKRQGARYVWNRT ELMQASLDPSVTHLMGLFE  
PGDMKYEIHRDSTLDPSLMEMTEAALRLLSRNPRGFFLFVEGGRIDHGHESRA  
YRALTETIMFDDAIERAGQLTSEEDTLSLVTADHSHVFSFGGYPLRGSSIFGLA  
PGKARDRKAYTVLLYGNGPGYVLKDGARPDVTESESGSPEYRQQSAVPLDEETH  
AGEDVAVFARGPQAHLVHGVQEQT FIAHVMAFAACLEPYTACDLAPPAGTTDAA  
HPG

(SEE ID NO: 3)

GGCAAGAGGGCGCGCTGGGAGCGCGCTGAGCGGGGAGAGGCGCTGCCCGACGGCCGGCCACAGGACCACCTCCCGGAG 79  
 AATAAGGGCCTCTTTATGGC M W L V T F L L L L D S L H K 15  
 ATG TGG CTG GTA ACT TTC CTC CTG CTC CTG GAC TCT TTA CAC AAA 143  
 A R P E D V G T S L Y F V N D S L Q Q V 35  
 GCC CGC CCT GAA GAT GTT GGC ACC AGC CTC TAC TTT GTA AAT GAC TCC TTG CAG CAG GTG 203  
 T F S S S V G V V V P C P A A G S P S A 55  
 ACC TTT TCC AGC TCC GTG GGG GTG GTG GTG CCC TGC CCG GCC GCG GGC TCC CCC AGC GCG 263  
 A L R W Y L A T G D D I Y D V P H I R H 75  
 GCC CTT CGA TGG TAC CTG GCC ACA GGG GAC GAC ATC TAC GAC GTG CCG CAC ATC CCG CAC 323  
 V H A N G T L Q L Y P F S P S A F N S F 95  
 GTC CAC GCC AAC GGG ACG CTG CAG CTC TAC CCC TTC TCC CCC TCC GCC TTC AAT AGC TTT 383  
 I H D N D Y F C T A E N A A G K I R S P 115  
 ATC CAC GAC AAT GAC TAC TTC TGC ACC GCG GAG AAC GCT GCC GGC AAG ATC CCG AGC CCC 443  
 N I R V K A V F R E P Y T V R V E D Q R 135  
 AAC ATC CGC GTC AAA GCA GTT TTC AGG GAA CCC TAC ACC GTC CCG GTG GAG GAT CAA AGG 503  
 S M R G N V A V F K C L I P S S V Q E Y 155  
 TCA ATG CGT GGC AAC GTG GCC GTC TTC AAG TGC CTC ATC CCC TCT TCA GTG CAG GAA TAT 563  
 V S V V S W E K D T V S I I P E N R F F 175  
 GTT AGC GTT GTA TCT TGG GAG AAA GAC ACA GTC TCC ATC ATC CCA GAA AAC AGG TTT TTT 623  
 I T Y H G G L Y I S D V Q K E D A L S T 195  
 ATT ACC TAC CAC GGC GGG CTG TAC ATC TCT GAC GTA CAG AAG GAG GAC GCC CTC TCC ACC 683  
 Y R C I T K H K Y S G E T R Q S N G A R 215  
 TAT CGC TGC ATC ACC AAG CAC AAG TAT AGC GGG GAG ACC CGG CAG AGC AAT GGG GCA CGC 743  
 L S V T D P A E S I P T I L D G F H S Q 235  
 CTC TCT GTG ACA GAC CCT GCT GAG TCG ATC CCC ACC ATC CTG GAT GGC TTC CAC TCC CAG 803  
 E V W A G H T V E L P C T A S G Y P I P 255  
 GAA GTG TGG GCC GGC CAC ACC GTG GAG CTG CCC TGC ACC GCC TCG GGC TAC CCT ATC CCC 863  
 A I R W L K D G R P L P A D S R W T K R 275  
 GCC ATC CGC TGG CTC AAG GAT GGC CGG CCC CTC CCG GCT GAC AGC CGC TGG ACC AAG CGC 923  
 I T G L T I S D L R T E D S G T V I C E 295  
 ATC ACA GGG CTG ACC ATC AGC GAC TTG CCG ACC GAG GAC AGC GGC ACC TAC ATT TGT GAG 983  
 V T N T F G S A E A T G I L M V I D P L 315  
 GTC ACC AAC ACC TTC GGT TCG GCA GAG GCC ACA GGC ATC CTC ATG GTC ATT GAT CCC CTT 1043  
 H V T L T P R K L K T G I G S T V I L S 335  
 CAT GTG ACC CTG ACA CCA AAG AAG CTG AAG ACC GGC ATT GGC AGC ACG GTC ATC CTC TCC 1103  
 C A L T G S P E F T I R W Y R N T E L V 355  
 TGT GCC CTG ACG GGC TCC CCA GAG TTC ACC ATC CGC TGG TAT CGC AAC ACG GAG CTG GTG 1163  
 L P D E A I S I R G L S N E T L L I T S 375  
 CTG CCT GAC GAG GCC ATC TCC ATC CGT GGG CTC AGC AAC GAG ACG CTG CTC ATC ACC TCG 1223  
 A Q K S H S G A Y Q C F A T R K A Q T A 395  
 GCC CAG AAG AGC CAT TCC GGG GCC TAC CAG TGC TTC GCT ACC CGC AAG GCC CAG ACC GCC 1283

FIG. 5

Q	D	F	A	I	I	A	L	E	D	G	T	P	R	I	V	S	S	F	S	415
CAG	GAC	TTT	GCC	ATC	ATT	GCA	CTT	GAG	GAT	GGC	ACG	CCC	CGC	ATC	GTC	TCG	TCC	TTC	AGC	1343
E	K	V	V	N	P	G	E	Q	F	S	L	M	C	A	A	K	G	A	P	435
GAG	AAG	GTG	GTC	AAC	CCC	GGG	GAG	CAG	TTC	TCA	CTG	ATG	TGT	GCG	GCC	AAG	GGC	GCC	CCG	1403
P	P	T	V	T	W	A	L	D	D	E	P	I	V	R	D	G	S	H	R	455
CCC	CCC	ACG	GTC	ACC	TGG	GCC	CTC	GAC	GAT	GAG	CCC	ATC	GTG	CGG	GAT	GGC	AGC	CAC	CGC	1463
T	N	Q	Y	T	M	S	D	G	T											465
ACC	AAC	CAG	TAC	ACC	ATG	TCG	GAC	GGC	ACC											1493

(SER ID NO: 5)  
(SER ID NO: 6)

FIG. 5

8f26 -----MMLVTFLLLLLSLHKARPED-----VGTSLYFVNDSLQQVTFSSS  
 D38492 --MKTPLLVSHELLLSLTSCLGFTWHRRYGHVSEEDKGFQPIFEQPIINTIYPEESLE  
 P20241EURO ---MWRQSTILAAALLVALLCAGSAESKGNRPPIRK-----QPAPGELLFXVAQONKESD  
 P32004EURA ---MVVALRYVWPLLLCSPCLLIQIPEEYEGHVMH-----PPVITEQSPR-RLVVFPTD  
 P35331G-CA -MMKEKSISASKASLVFPLCQMHISALDVLPLDSKLLLELS-QPPTITQOSP-K-DYIVDPRE  
 Q02246XONI -MGTATRRKPHLLLVAALVALVSSSAWSSALGSQTT-----FGPVFEDQPLSVLPFEESTZ  
 U11031 -----MLSWKQLILLSFIGCLAGELLL-----Q-----QPVFVKEPSNSIFPVGSED  
 X65224 MVLHSHQTYAGIAPALCLHLHLSAIEVPLDSNIQSELP-QPPTITKQSVK-DYIVDPRE

8f26 VGVVVPCPAAGSPSAALRWYLATGDDIYDVPHIRHVHANG--TLQLYPPSPSAFNSFIHD  
 D38492 GKVSINCRARASPPFVYKWRMN-NQDVDLTN-DRYSMV----GQNLVINNFDPKQK-D--A  
 P20241EURO NPFTIECEADQPEPEYSWIKN-GKFPDQWQAYDNRLRQPG-RGTLVITIPKDED----R  
 P32004EURA D-ISLKCEASGKPEVQPRWTD-QVHFKPEELQVTVYQSPHSQSFTITGNNSNFAQRFO  
 P35331G-CA N-IVIQCEAKGKPPPSFSWTRN-GTHFDIDKDAQVTMKPN--SGTLVVNINMGVKAAYE  
 Q02246XONI EQVLLACRARASPPATYRWQON-GTEMKLEPQSAHQLV----GQNLVINNPTKAQ-D--A  
 U11031 KKITLNCARGNPSPHYRWQLN-GSDIDTSLDHRYKLN----GQNLVINPNRNW-D--T  
 X65224 N-IFIECEAKGNPVPTFSWTRN-GKFFNVAKDPKVSMMRR--SGTLVIDFHGGGRPDDE

8f26 NDYFCTAENAAGKIRSPNIRVKAUFREPYTVRVEDQQRSMR-GNVAVFKCLIPSSVQEVVS  
 D38492 GITYCLASNNYGMVRSTEATLSFGYLDPPFPEDRPEVKVKEGKGMVLLCDPPYHFPDD-L  
 P20241EURO GHYQCFASNEFGTATSNSVYVRKAELNAFKDEAAKTLEAVEGEPFMLKCAAPDGFPS--P  
 P32004EURA GIYRCFASNKLGATMSHEIRLMAEGAPKWPKEVTVKPVVEEGESVVLPCNPPPSAEP--L  
 P35331G-CA GVYQCTARNERGAATSNIVIRPVSPLWTKEKLEPNHVREGDSLVLNCRFPVGLPP--P  
 Q02246XONI GVYQCLASNVGTVVSREAILRFGFLQFESKEERDFVKAHEGWGVMLPCNPPAHYPG--L  
 U11031 GSTQCFATNSLQTVSREAKLQFAYLENFKSRMRSRVSVREGQGVVLLCGPPPHSGE--L  
 X65224 GETQCFARNDYGTALSSKIHQLQVRSPLWPKKVDVIEVDEGAPLSLQCNPPPGLP--P

8f26 VVSWEKDTVSIIE-----NR--FFITYHGGLYISDVQKED--ALSTYRCITKHYSGET  
 D38492 SYRWLLNEFPVFTIM---DKRRFVSQ-TGNLYIANVESSD---RQNTSCFVSS--PSIT  
 P20241EURO TVNMNIQESIDGSIKSINNSR--MTLDPEGNLWFSNVTREDASSDFYACSATSVFRSEY  
 P32004EURA RIYWNKSKILHIQ-----DER--VTMQQNGNLYTANVLTSDN--HSDYICHAHFQTRTI  
 P35331G-CA IIFWMDNAFQRLPQ-----SER--VSQQLNGDLYFSNVQPEDT--RVDYICYARFNHTQTI  
 Q02246XONI SYRWLLNEFPNFIPT---DGRHFVSQ-TTGNLYIARTNASD---LQNTSCLATSHMDFST  
 U11031 SYAWVTFNEYPSFVEE---DSRRFVSQ-ETGHLIYAKVEPSD---VQNTCTCVTS--TVTN  
 X65224 VIFWSSSMETPIHQ---DKR--VSQCGNGDLYFSNVMLQDA---QTDYSCNARFHTHTI

8f26 RQSNGARLSVTDPAES-----IPTILDGFHSQEV---WAGHTVEL  
 D38492 KSVFSKFIPLIPERTT-----KPYPADIVVQFKDIY--TMMQONVTL  
 P20241EURO KIGNKVLLDVKQMGVSASQ-----NKHPPVROYVSRQOS-LALRGKRMEL  
 P32004EURA IQKEPIDLRVKATNSMID-----RKPRLLFPTNSSSHLVALQCGQPLVL  
 P35331G-CA QQKQPISVKVFSTKP-----VTERPPVLLTPMGSTSNKVELRGNVLLL  
 Q02246XONI KSVFSKFAQLNLAAEDTR-----LFAPSIKARFPAETY--ALVGQQVTL  
 U11031 ARVLGSPTPLVLRSDGVMG-----EYEPKIELQFPETLP--AAKGSTVKL  
 X65224 QQKNPYTLKVTKKPHNETSLRNHTDMYSARGVTETTPSFMPYGTSSSQMVLRGVDLL

8f26 PCTASGYPIPAIRWLKDRP--LPADSRWTKRITGLTISDLRTEDSGTYICEVTNTFGSA  
 D38492 ECFALGNFVPDIRWKVLEP--MPTTAEISTSGAVLKIFNIQLEDEGLYECEAENIRGKD  
 P20241EURO FCIYGGTPLPQTWVSKDGQRIQWSDRITQGHYKSLVIRQTNFDDAGTTCYDVSNVGNA  
 P32004EURA ECIAEGFPTPTIKWLRPSGPM-PADRVTYQNHNTLQLLKVGEEDDGEYRCLAENSLGSA  
 P35331G-CA ECIAAGLPTFVIRWIKEGGEL-PANRTFFENFKKTLKIIDVSEADSGNYKCTARNTLCST

FIG. 6



Q02246XONI ECFAGNFVPRIKWRKVDG----SLSPQWTTAEPTLOIPSVSFEDQTYECLAENSKGRD  
U11031 ECFALGNFVPQINWRRSDQMP--PPTKIKLRKPNGLZIPNFQQEDTGSYECIAENSRGKN  
X65224 ECIASGVFAPDIDWYKKGEL-PAGKTKLENFNKALRISNVSEEDSGEYFCIASNMKMSI  
\* \* \* \*

8f26 E-ATGILMVIDPLHVTLTTPKRLKTGIGSTVILSCALTGSPEPTIRWYRNT-----  
D38492 K-HQARIYVQAFPEWVEHINDTEVDIGSDLYWPGVATGKPIPTIRMLKNG-----  
P20241EURO QSFSIILNVNSVPYFTKEPEIATAAEDDEVVFECAAGVPEPKISWIHNGKPIEQSTPNP  
P32004EURA R-HAYYVTVAAAPYWLHHPQSHLYGPGETARLDCQVQGRPQPEVTWRINGIPVEELAKDQ  
P35331G-CA H-HVISVTVKAAPYWTAPRNLVLSPGEDGTLCRANGNPKPSISWLTNGVPIAIAPEDP  
Q02246XONI T-VQGRITVQAQPEWLKVISDTEADIGSNLRWGCAAAGKPRPTVRMLRNGEPLASQNR--  
U11031 V-ARGRLTYAKPYWVQLLKDVTAVEDSLYWECRASGKPKPSYRMLKNGDALVLEER--  
X65224 R-HTISVRVKAAPYWLDEPQNLILAPGEDGRLVCRANGNPKPSIQWLNVNGEPIEGSPPNP  
\* \* \*

8f26 -----E-----LVLFDRAISIRGLSN-----  
D38492 -YAYHKQELRLYDVTPEFENAGMYQCIAENAYGTIYANAELKILALAPTFFEMNPMKKKILAA  
P20241EURO RRTVTDNTIRIINLVKGDTCNGYGCNATNSLGYYVYKDVYLVNQAEPP--TISEAPAAVSTV  
P32004EURA KYRIQRGALILSNVQPSDTMTVTCENNRHGLLLANAYTYVVLPA-KILTADNQTMAV  
P35331G-CA SRKVDGDTIIFSAVQERSAAYVQCNAESNEYGYLLANAFVNVLAEP--RILTPANKLYQVI  
Q02246XONI -VEVLGDLRF SKLSLEDSCMYQCAENKHGTIYASAEALVQALAPDFRLNPVRRLIPAA  
U11031 -IQIENGALTIANLVSDSQMFQCIAENKHGLIYSSAELKULASAPDFSRNPMQMIQVQ  
X65224 SREVAGDTIVFRDTQIGSSAVYQCNASNEHGYLLANAFVSVLDVFP--RILAPRNQLIKVI

8f26 -----ETLLITSAQKSHSGAYQCPA  
D38492 KGGRVIIIECKPKAAPKPKFSWSKGTWLVNSSRILIWED-GSLEINNITRNDGGIYTCFA  
P20241EURO DGRNVTIKRVNGSPKPLVKWLRASNWLT--GGRYNVQANGDLEIQDVTFSAGKYTCYA  
P32004EURA QGSTAYLLCAGAPVPSVQWLEDGTTVLQDERFFPYANGTLGIRDQANDTORIYFCLA  
P35331G-CA ADSPALIDCAYFGSPKPEIEWFRGVKGSILRGNEYVFDHNGTLEIPVAQKSTGTYTCVA  
Q02246XONI RGGEILIPCQPRAPKAVVLWSKGTIILVNSSRVTVTPD-GTLIIRNISRDEGKYTCFA  
U11031 VGSVLIDCKPSASPRALSFWKKGDTVVREQARISLLND-GGLKIMNVTKADAGIYTCIA  
X65224 QYNRTRLDCCPFFGSP IPTLRWFKNQGNMLDGNGYKAHENGSLMSMARKEDQGIYTCVA  
\* \* \*

8f26 TRKAQTAQDFAIIALEDGTPRIVSSFSEKVVNPGEQFSLMCAAKGAP--PFTVTWALDDE  
D38492 ENNRGKANSTGTLVITNPT-RIILAPINADITVGENATMQCAASFDPSLDLTFVMSFNGY  
P20241EURO QNKFGIQAQDQSLVVKHT-RITQEPQNYEVAAGQSATFRCEAHDDTLEIEIDWWDGQ  
P32004EURA ANDQNNVTIMANLKVKDAT-QITQGPRTSTIEKKGSRTVFTQASFPDPSLQPSITWRGDGR  
P35331G-CA RNKLGKTQNEVQLEVKDPT-MIIKQPQYKVTORSQAQASPECVVKHDPTLIPTVTWLKD--  
Q02246XONI ENFMGKANSTGILSVRDAT-KITLAPSSADINLGDNLTLQCHASHDPTMDLTFMTLDDF  
U11031 ENQFGKANGTTQLVVTETPT-RIILAPSNDVAVGESIILPCQVQHDPLLDIMFAMWFNGT  
X65224 TNILGKVEAQVRLEVKDPT-RIVRGPEQVVKRGSMPRLHCRVKHDPTLKLTVTWLKD--  
\* \* \*

8f26 PIVRDGSHRTNQYMS----- (SEE ID NO: 7)  
D38492 VIDFNKEITNIHYQRNFMLDANGELLIRNAQLKHAGRYTCTAQTIVDNSSASADLVVRGP ( " 8)  
P20241EURO SIDFEAQPR----FVKTNNDN--SLTIAKTMELDSGEYTCVARTRLDEATARANLIVQDV ( " 9)  
P32004EURA --DLQELGD---SDKYFIEDG--RLVIHSLDYSQGNYSCEVASTELDVSRAQLLVGS ( " 10)  
P35331G-CA --NNELPDD---ERFLVGKD--NLTIMNVTDKDDGTYTCIVNTTLDVSASAVLTVVAA ( " 11)  
Q02246XONI PIDFDKPGG--HYRRTNVKETIGDLTILNAQLRHGGKYTCMAQTUVDSASKEATVLVRGP ( " 12)  
U11031 LTDFKKDGS--HFEKVGGSSS-GDLMIIRNIQLKHSGKYVCMVQTVGDSVSSAAELIVRGS ( " 13)  
X65224 --DAPLYIG----NRMKKEDD--GLTIYGVAEKDQCDYTCVASTELDKDSAKAYLTVLAI ( " 14)

FIG. 6

## INTERNATIONAL SEARCH REPORT

International application No.  
PCT/US97/20201

## A. CLASSIFICATION OF SUBJECT MATTER

IPC(6) : C07H 21/04; C07K 14/47; C12N 5/16, 15/70, 15/79; C12Q 1/68

US CL : 435/6, 320.1, 325; 530/350; 536/23.5

According to International Patent Classification (IPC) or to both national classification and IPC

## B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

U.S. : 435/6, 172.3, 320.1, 325, 365; 530/350; 536/23.1, 23.5; 935/22, 24, 27, 79

Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practicable, search terms used)

APS, STN (Biosis, CAPlus, LifeSci, Medline, INPADOC, WPIDS), Genbank, EMBL, Pir

## C. DOCUMENTS CONSIDERED TO BE RELEVANT

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
A	US, 5,525,486 A (HONJO et al.) 11 June 1996, see entire document.	1, 3, 4
A	US, 5,536,637 A (K. JACOBS) 16 July 1996, see entire document.	1, 3, 4

☐

Further documents are listed in the continuation of Box C.

☐

See patent family annex.

* Special categories of cited documents:	*T* later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
*A* document defining the general state of the art which is not considered to be of particular relevance	*X* document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone
*E* earlier document published on or after the international filing date	*Y* document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art
*L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)	*G* document member of the same patent family
*O* document referring to an oral disclosure, use, exhibition or other means	
*P* document published prior to the international filing date but later than the priority date claimed	

Date of the actual completion of the international search

27 JANUARY 1998

Date of mailing of the international search report

23 FEB 1998

Name and mailing address of the ISA/US  
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Box PCT  
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## ptrAP3



FIG. 1

## ptrAP3 vector sequence

AAGCTTGGCTGTGGAATGTGTGTCAGTTAGGGTGTGGAAAGTCCCCAGGCTCCCCAGCAGGCAGAAAGTATGC  
 AAAGCATGCATCTCAATTAGTCAGCAACCAGGTGTGGAAAGTCCCCAGGCTCCCCAGCAGGCAGAAAGTATGC  
 AAAGCATGCATCTCAATTAGTCAGCAACCATAGTCCCGCCCTAACTCCGCCCATCCCCCCCCCTAACTCCGC  
 CCAGTTCGGCCCATTTCTCCGCCCCATGCTGCTGACTAATTTTMTTATTTATGCAAGAGGCCGAGGCCGCTCGG  
 CTTCTGAGCTATTCAGAAAGTAGTGAGGAGGCTTTTMTGGAGGCTTAGGCTTTTGCAAAAAGCTCCTCCGAT  
 CGAGGGGCTGGCATCTCTCTTCAAGGCCCTCCGCCCTAAGCTTAGGCGGCCCATCCACGCCCGTTGAGTCCG  
 GTCTGCGGCCCTCCGCCCTGTGGTGCCTCTGAACTGGCGTCCGCCCTTAGGTAAGTTTAAAGCTCAGGTCG  
 AGACCGGGCTTTTGTCCGCCCTCCCTTGAGGCTTACCTAGACTCAGCGGCTCTCCAGGCTTTGCGCTGACC  
 CTGCTTGCTCAACTCTACCTCTTGTCTTCTGCTTTCTGCTGCGCTTACAGATCCAAAGCTCTGAAAAACC  
 AGAAAGTTAACTGGTAAGTTTACTCTTTTGTCTTTTATTTAGGTCGCCAGGTCGCCGATCCGGTGTATCCAA  
 ATCTAAGAAGTCTCTCTCAGTGAGTGTGTGCTTTACTTCTAGGCTGTACGGAAGTGTACTTCTGCTCTAA  
 AAGCTCGGGAATTCGACACACCGTAGTCTTTTACCGCCGCTGAGCGCTCCAGCCGACCTACA  
 AGCGCTCTATATGATGAGTGTACGCGGACGAGGACTTCTTGAAGCAGGCCAAGCGAGCGCT  
 CGCGGAGTTTGCCTACGGAAGCGGCATAGGACATGTTGCGCTTCCCGCTGACGAGGCG  
 AAGCCAAACAGCTAGGCTTAAAGCGCGTGACACTGACAGCAGGTGCTGCGCAGCTTTCACCGT  
 CCGAAGAAAGAGCGCGCTTAAAGCGCGAGTCTGCTGACTTGGCAGCCAGCTTGCAGCTGAT  
 GGTACCCCAAGCGCCAGGGAAGTGAAGATGCTCTTGGAAAAATGACCGTGGAGCGCTGCGCTG  
 GAGCCCGAGGTCGCGGTGCGGCGAATCAAGCAAGTGGCAGCGGAGCTGGCGCTGACAGACCG  
 TGGACGTTTCAGATAGCCAGCCAGGAGTACACTAGTATTTGCCACTGCGACAGAGGCGCATGGA  
 GACACAAACGTCGCCCGCTTCCCTAAGCTCAGATCATCTTACTGAGGAGGAGAAACCGGACTTCTG  
 GAACCGTTAGGCAGCTGAGGCTTGGCTGCGGCAAGTACCTTGAAGCTGACAGACAGCGGCCAAGTACCT  
 CATCATCTTCTGCTGCTGATGCGATGCGGCTGCTTACGCTGACAGCTGCTGAGGATGCTTAAAGCGCGAGAG  
 GGACAAACTGGCGCTTGAATATCTCTGCGCATGAGACCGCTTGCATATGCGGCTCTGCTTCAAGACATACAA  
 TGTAGACAAACATGCTGACAGAGTGGAGGACAGCGGCAAGGCTTACTTGGCGGCTCAAGCGCAAGTTCGA  
 GACCATGCGCTTACAGTGGAGCGCGGCTTTAAGCAAGTGCACACGACAGCGGCAAGTACAGTCACTGCT  
 GATGAATCGCGGCAAGAAAGCAAGTAACTGCTGCTGCTGCTTAAACACCAAGCAAGTTCAGCAGCGCTGCG  
 AGCGGCGACCTTACGCGGCAAGCTTAAAGCGCAAGTCTTACTGAGGAGGCGAAGTGGCTGCGCTGCGCGGCA  
 GCGCGGCTGCGCAGGACATGCTTACGCGAGCTCATCTGCAACATGCAATGCAAGTCAAGTGGTGGAGCGG

FIG. 2



TTATGGCAGCCTGCATRAATTCTCTTACTGTCAAGCCATCCGTAAGATGCTTTTCTGTGACTGGTGAGTACT  
CAACCAAGTCATTCTGAGAAATACTGTATGCGGCGACCGAATTGCTCTTGCCCGCGGTCAATACGGGATAATA  
CCGCGCCACATAGCAGAACTTTAAAGTGCTCATCATTTGGAAAACGTTCTTCGGGGCGAAAACCTCTCAAGGA  
TCTTACCGCTGTTTAGATCCAGTTCGATGTAACCCACTCGTGACCCCAACTGATCTTCAGCATCTTTTACTT  
TCACCAGCGTTTCTGGGTGAGCAAAAACAGGAAGGCAAAATGCCGCAAAAAGGGAATAAGGGCGACACGGA  
AATGTTGAATACTCATACTCTTCCTTTTTCAATATTATTGAAGCATTATCAGGCTTATTGTCTCATGAGCG  
GATACATATTTGAATGTATTTAGAAAAATAAACAAATAGGGGTTCCCGCCACATTTCGCCGAAAAGTCCAC  
CTGC

(SEE SA 100)

2

FIG. 2

FIG. 3

MLLLLLLLGLRLQLSLGIIPVEEENPDFWNREAAEALGAACKKLQPAQTAACKNLI  
IFLGDMGMGVSTVTAARILKGQKKDKLGPFIPLAMDRFPYVALSKTYNVDKHVDP  
SGATATAYLCGVKGNEFQTIGLSAAARFNQCNTTRGNEVISVMNRAKKAGKSVG  
VTTTRVQHASPAGTYAHTVNRNWYSDADVPA SARQEGCQDIATQLISNMDIDVI  
LGGGRKYMFRMGTPDPEYPDDYSQGGTRLDGKNLVQEWLAKRQGARYVWNRT  
ELMQASLDPSVTHLMGLFEPGDMKYEIHRDSTLDPSLMEMTEAALRLLSRNPRGFF  
LFVEGGRIDHGHESRAYRALTETIMFDDAIERAGQLTSEEDTSLSVTADHSHV  
FSFGGYPLRGSSIFGLAPGKARDRKAYTVLLYGNGPGYVLKDGARPDVTESESG  
SPEYRQQSAVPLDEETHAGEDVAVFARGPQAHLVHGVQEQT FIAHVMAFAACLE  
PYTACDLAPPAGTTDAAHPGRSVVPEALLPLAGTLLLLLETTATAP

(SEE IN SEQ. 2)

FIG. 4

II PVEEENPDFWNREAAEALGAACKKLQPAQTAACKNLIIFLGDMGMGVSTVTAARI  
LKGQKKDKLGPFIPLAMDRFPYVALSKTYNVDKHVDP SGATATAYLCGVKGNEFQ  
TIGLSAAARFNQCNTTRGNEVISVMNRAKKAGKSVG VTTTRVQHASPAGTYAH  
TVNRNWYSDADVPA SARQEGCQDIATQLISNMDIDVILGGGRKYMFRMGTPDPE  
YPDDYSQGGTRLDGKNLVQEWLAKRQGARYVWNRT ELMQASLDPSVTHLMGLFE  
PGDMKYEIHRDSTLDPSLMEMTEAALRLLSRNPRGFFLFVEGGRIDHGHESRA  
YRALTETIMFDDAIERAGQLTSEEDTSLSVTADHSHVFSFGGYPLRGSSIFGLA  
PGKARDRKAYTVLLYGNGPGYVLKDGARPDVTESESGSPEYRQQSAVPLDEETH  
AGEDVAVFARGPQAHLVHGVQEQT FIAHVMAFAACLEPYTACDLAPPAGTTDAA  
HPG

(SEE IN SEQ. 3)





O	D	P	A	I	I	A	L	E	D	C	T	P	R	I	V	E	S	P	S	418
CAG	GAC	TTT	GCC	ATC	ATT	GCA	CTT	GAG	GAT	GGC	ACG	CCC	CGC	ATC	GTC	TGC	TCC	ATC	AGC	1343
E	K	V	V	N	P	Q	E	Q	F	S	L	M	C	A	A	K	G	A	P	435
GAG	AAG	GTG	GTC	AAC	CCC	GGC	GAG	CAG	TTC	TCA	CTG	ATG	TGT	GGG	GCC	AAG	GGC	GCC	CCG	1401
F	P	T	V	T	H	A	L	D	D	E	P	I	V	E	D	D	S	H	R	455
CCC	CCC	AGG	GTC	ACG	TGG	GCC	CTC	GAC	GAT	GAG	CCC	ATC	GTC	CGG	GAT	GGC	AGC	CAC	CCC	1483
T	N	Q	Y	F	M	S	D	C	T											855
ACC	AAC	CAG	TAC	ACC	ATG	TCC	QAC	GGC	ACC											1491

(SER 18 NOV: 5)  
(SER 18 NOV: 6)

FIG. 5

8126  
 D38492  
 P20241EURA  
 P32004EURA  
 P35331G-CA  
 Q02246XONI  
 U11031  
 X65224  
 -----MHWLVTFLLLLDSLKKARPE-----VGTSLVFWHDSLQGVTFSS  
 --MHTYLLVSKLLLSLSCLOSTWHRAYUNGVSSEKRGTCPIFEQPIINTIYPKESE  
 ---MWRQSTYLLAALLVALLCAGRAESKQTHPPFLIT-----QPAPGELLFKVRAQONKESD  
 ---MVALAYVWFLLOSFCLLIQIPREYEGHEVME-----FVUTEQSPR-ELWVFTD  
 -MCKKCHISAKASLVVFLCQIEALOVPLDSKLLSELS-QPFTITQGSFK-DYIVDFRE  
 -MPTATKXPHLLLVAAVALVSEBANSALGSETT-----FQVVEDQPLSVLPFEESTZ  
 -----MLNKOLILLSFIQCLAGELLL-----Q-----QFUTVKEPENSIPFVGSZD  
 MULHSHQLTYAQIAFALCLMHLISAIEVPLDSNTQSELP-QPFTITRQSVK-DYIVDFRD

8126  
 D38492  
 P20241EURA  
 P32004EURA  
 P35331G-CA  
 Q02246XONI  
 U11031  
 X65224  
 VGVVVFQBAAGSPSAALRWYLATGDDIYDVPHIRVHANG--TLQLYTFSPFAVHSPIND  
 GKVSLNGARASFPFUYKQNN--MGOVDLTH-DRYEMV---GGLVIVNTPKQK-D--A  
 NPFTIEGADQGFPEYFWING--GQVFDQAYDNKELRQFG-RQTLVITIKHED-----R  
 D-TSLKGEASQKPEVDFPMTD--GVHFKPKSELQVTVYQSPHGEFTITOMNENPAQRPO  
 N-IVIQCEAKQKPPPEFWTRN--GTHFDIDKDAQVTCKFH--SGLVUNINRQVKAAYE  
 ECVLLACARASPPATYKWTGN--GTEMKLEPGJRHQLV---GGLVIVNTPKQK-D--A  
 KKITLNCZARGNPSPHYRWQIN--QSDIOTSLDNVYKLN---GGLVIVNTPKQK-D--T  
 N-IFTECEAKQKPPVTFSTWTRN--GKTFNVARDPKVENDRA--SGLVIVDFHGGKRPDDYE

8126  
 D38492  
 P20241EURA  
 P32004EURA  
 P35331G-CA  
 Q02246XONI  
 U11031  
 X65224  
 NDYFCTAEDAAOKIRSPMIXVKAFTREPYTVREDORENR--GVAVVFKCLIPSSVQETVS  
 GITYCLASNNYQNVSTKATLSFGTLQFPFPEDRPEVYVUEGKXGVLLDPFYKFFDE-L  
 GHTQCFASNEPGTATNSVYVRAELNAPIDEAAKTLAEGEFPFKLCAALPOFFPS--P  
 GITYCFASNELOTAMSHIILMAHAPKHTKTVKPVVEDEGESEVLSCHFPPEABP--L  
 GVTQCTARNEQAALINNVIRPSPKPLWTKHLEPHVABEDDELVLNCPFFVLFP--P  
 GVTQCLASNPVQTVVSKLALLPQPLQEFSEKEREPUKAREGQVNLKCHFPARYG--L  
 GSTQCTATNELOTIVERDAKLOFAYLKNYKEMERVSUREGQGVLLQOPFPKSGE--L  
 GETQCTANDYOTALSSKIMLQVERSPIMPKERVVIEUDEGAPLELQGNFPFGLFP--P

8126  
 D38492  
 P20241EURA  
 P32004EURA  
 P35331G-CA  
 Q02246XONI  
 U11031  
 X65224  
 VVSWKMDTVSIIPE-----NR--FFITYHGGLYISOVQKED--ALSTREITTHKYSGET  
 SYWLLNEFPVITM---DNKRFVSC--TGNLYIANVERED---KONTSCPVES--PSIT  
 TVNHTIGESIDGSIKSIKNER--MTLDPSENLWFMVTRDASSDFFYACSATSVTRBY  
 RYNSHKKILHINO-----DER--VTNQGNGNLYTAMVLTSDN--HSDYICHANFPOTRTI  
 IIPQENAFQRLFPQ-----SER--VSGQKNGDLXFNVOPEET--RVDYICYARFHTQTI  
 SYWLLNEFPVITM---DGRHVFSC--TGNLYIARTNAED---LONTSEGLATERDOPST  
 SYWVTHNYFPSTPE---DSKRFVSC--STGHLYIAKVEPED---VONTSEVTS--TVTR  
 VTPWHSSEHPIHQ---DNK--VSGQKNGDLXFNVOPEET--QTDVSENAFPKTHTI

8126  
 D38492  
 P20241EURA  
 P32004EURA  
 P35331G-CA  
 Q02246XONI  
 U11031  
 X65224  
 PQSNGARLSVTHPAES-----IFTILDGTHSQEV---WAGHTVEL  
 KSVFSEKPIPLIPERTT-----KPYFADIVVQFXDIY---TMOGQNVTL  
 KICKVLLDQVQKQVVSAGQ-----NKHSEVRQYVSRQS-LALAGKMKEL  
 IQKEPIDLRVKTATNEMID-----RKPALLFPNSESSELVALQGGFLVL  
 QKQKQPISVKVFSTKP-----VTERFPVLLTFMSTENKVELRGNVLL  
 KSVFSEPAQNLAAEDTA-----LFAPSIRAKFPARTV---ALVGOQVTL  
 ARVLGSPPTFLVLRSDGVK-----EYEPKIELQPFETLP---AAKGSTVKL  
 QKQKQPIVTVKXENPHDETSLRNHTNMYBARGVTETTFEFMYFYGTSSSQMVLQGVLL

8126  
 D38492  
 P20241EURA  
 P32004EURA  
 P35331G-CA  
 PCTASGYFIPAIRMLKDCRP--LPADSRMTKUTGLTISDLRTSDGTTCVNTFQSA  
 ECFALGNPVPIKMKVLEP--MPTTARISTEGAVLKIFNIQLEDEGLTECAENINGKD  
 PCIVGCTPLPQTVMKQGGRIQWDRITCGHYGKSLVZAGTNPDEASTTTCVSNQVQNA  
 SCIAAGSPPTFTIWLRFSGPM--PADRVYQNTENTLQLLKVEEEDGCTRCALANISLOSA  
 SCIAAGLPTFVIRWIKEGEL--PANRTTFTENPKTKIIEVSEADSCNTTCTARTLIGST

FIG. 6

Q02246XONT  
U11031  
X65224

BETAFGNFVPRHXKXVDG----SLSPQNTTAEPTLQIPSVSPEDGPTTCTEAEHSEKRD  
BGFALGNFVPGLEWRKSDQMP--PFTKIKLARFNGVLEIPSTQQEDTGSTFCTAEHSEKRD  
BCLAGVBPAPDINWYKKGEL--PAGKTELENFNKALRISNVSEEDSGSTFCLAHNGQSI  
\* \* \* \*

8226  
D38492  
P20241EURO  
P32004EURA  
P35331G-CA  
Q02246XONT  
U11031  
X65224

R-ATGILKVIDPLHVTATPCKLXTGIGSTVILSCALTCSEPTTINWYANT-----  
K-HQARIYVQAFPEWVEHINDTENDIOGDLVWPGVATGCRIPITIKWLEKQ-----  
QSF8IILNUNSVPTTKEPELATAAEEVVEDEAAQVPEPLISWTHNGKPIEQSTFEP  
R-HAYYVTVAAAPYWLKXQSHLYGPGSTARLDQVQGRQPELWAIKQIPVEELAKDQ  
H-BVISVTVMARPYWITAPRNLVLSPOEDTLICRAMEKPKS;EWLTKRVTIATAPEDP  
T-VGGKTIYQAGPEWLVSTSTEDADIGENLWQCAAGKPAFTVWLEKQEZLASQGR--  
V-ARGKLTYYAKPYWVQLKDVETAVDESLYWCERAGKPKPSYRWLKNQDALVLSER--  
R-HTISVRVKAAPYWLDEPQNLILABGEDQRLVCRANGNPKPSICWLVNKEPIEGEPNP  
\* \* \* \*

8226  
D38492  
P20241EURO  
P32004EURA  
P35331G-CA  
Q02246XONT  
U11031  
X65224

-----8-----LVLFDKATSIKGLSH-----  
-YAYHKQELRLYDVTENAGHYQCIENAYQTIYANAEIKILALAPTTEKRMWKKILAA  
RRTVTENTIRIINLVKQNTGNYOCNATNSLGYVYKDVVLNVQASBP--TISKAPAAVSTV  
KYRIQKQALILSNVQPSDTHVTCERANNHGLLANAYTYVQLPA-KILTADNQTMAV  
SHKVGDTIIFSAVDERSAUVYCCNASNEYGYLLANAFVWMLASBP-KILTANLTYQVT  
-VEVLADLRFENLSLEDSECHYQCVANNGOTIYASALLAVQALAPDFALNPURRIPAA  
-IQIENGALTIANLVSDGHTQCIENHNGLYSSALVULASAPDFENHNGHNIQVQ  
SREVAGDTIVFRDTCIGENAYVQCAENENGYLLANAFVSVLDVTF-RILAPRQGLIKVI

8226  
D38492  
P20241EURO  
P32004EURA  
P35331G-CA  
Q02246XONT  
U11031  
X65224

-----ETLLITEAQKESGAYQCPA  
KQGRVJIECKEKAAPKPKFSWSKGTENLVNEERILINED-GELZINXITRNDGIIITEFA  
DGRVYTIKGRVNGSPKPLVWLRASWLT--GGRYNVQANGDLEIQDVTPEAGETCYA  
QOSTAYLLCKAFQAFVFEVQWLDGDTTLQDERFFPYANGTLGIRDLQANDTCRTYELA  
ADSPALIDCAYTCSEKPEIENWFGVKGSLRQNEYVTHQNTLEIPVAGKDTGTTTCVA  
RQGEIILIPQCPRAAPKAVVLWSKDTIELVNSERVVTTPD-UTLIINISRSDEGNYTEFA  
VGLVILDCXKPSASPPALSFWXKQDTVREQARISLND-GGLKINWYTKDAGIYTOIA  
QYNTRELCBPFFC29IYTLNFKNGQGNLQOCHYKAKENGSLNEMARKEDQGLYTCVA  
\* \* \* \*

8226  
D38492  
P20241EURO  
P32004EURA  
P35331G-CA  
Q02246XONT  
U11031  
X65224

TRKQTAGDFAIALLEDOTPRIVSESPSEKVNPOEQFSLMCAKNGAP--PFTTQOLDDE  
ENHGTAFNSTOTLVITNPT-RILAFINADITYGRNATEQCAAFDPDELDTVWENQY  
QWKFGEIQADOSLVKENT-RITQSPQNYEVAAGQSATFCEZANDOTLEIEIDWKKDQ  
ANDQENVTIMASLVKDAT-CITQGPSTIEKGRSVTFTQASTDFELQPSITMRGDR  
RNELGKTKQNEVQLAVKDP-MIIGKPSYKVTQELAGASFEVIXHDTLIPITVWLKD--  
ENFMKANSTUILSVRDAT-KITLAPSSADINLGENLTLOCKAKHDTMDLTFTMTLDDP  
ENQFORANGTQLVUTEPT-RILAPSKMDVAVGESIILPCQVQKDFLLDINFAMYTGT  
TWILKVBQAQVLEVKDPT-RIVRGEDQVVKRQSKPRLNCRVMDPTLKLITVWLKD--  
\* \* \*

8226  
D38492  
P20241EURO  
P32004EURA  
P35331G-CA  
Q02246XONT  
U11031  
X65224

FIVEDGSHRTNQYTMS----- (SEQ ID NO: 1)  
VIDFNKEITNHYQGNFMDANGELLIRNAQLNAGNYTCTAQTIVNNSASADLVVRGP ( " 2)  
SIDFHAQPR----FVXTNDN--SLTIKTHELOSCQYTCVARTLNEATHRAMLIVQOV ( " 3)  
--DLSLOD---SPRYFLEDG--RLVIHELDPYSGQNTFCVASTELDUVESRAQLVVRG ( " 4)  
--NRLEEDD---ERFLVQKD--WLTIMVTDKDDGTYTCTIVTTLDEVSAGAVLTVAA ( " 5)  
PIDFDRPGG--HYRATNVCTIGDLTILNAQLNAGNYTCTAQTIVNNSASADLVVRGP ( " 6)  
LTDYKKGDS--HFEKVGGES--QDLNIRFIQLKESGNTVCHVQTOVDVSSAELIVRG ( " 7)  
--DAPLYIG----NRKQEDD--GLTIYGVASKDGGTYTCVASTELDKDSARAYLTVLAI ( " 8)

FIG. 6

